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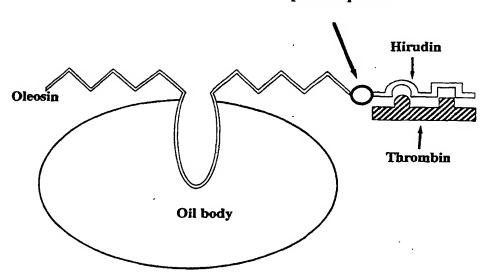
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(54) Title: OIL BODIES AND ASSOCIATED PROTEINS AS AFFINITY MATRICES

Spacer sequence



(57) Abstract

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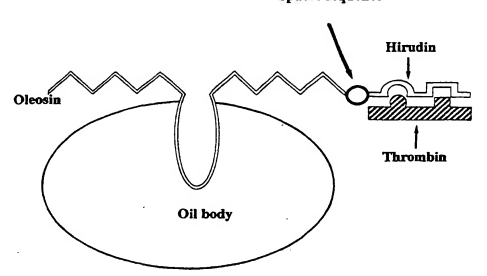
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TITLE: Oil Bodies and Associated Proteins as Affinity Matrices FIELD OF THE INVENTION

This invention relates to the use of oil bodies and their associated proteins as affinity matrices for the separation and purification of target molecules from samples.

BACKGROUND OF THE INVENTION

Within the general field of biotechnology, the ability to effectively separate and purify molecules from complex sources, such as living cells, blood serum, or fermentation broth, is of critical importance. Applications in industry and research laboratories (where, for example, purified or partly purified proteins are used) are numerous and well documented in prior literature. See, for example, R. Meadon and G. Walsh in *Biotechnological Advances* 1994, **12**: pp 635-646.

The majority of currently employed techniques for the separation of molecules capitalizes on the innate physical and chemical properties of the molecule of interest. Affinity-based purification technologies are unique in that they exploit the highly specific biological recognition between two molecular species which form an affinity pair. Binding of the two entities of the affinity pair occurs in almost all instances as a result of relatively weak chemical interactions, known as non-covalent bonds. Some art-recognized and commonly used affinity pairs include antibodies and their binding antigenic substances, nucleic acid binding proteins and nucleic acids, lipid binding proteins and lipids, lectins and carbohydrates, streptavidin/biotin complexes, protein A/immunoglobulin G complexes, and receptors and their binding molecules.

In general, affinity-based purification processes require that one member of the affinity pair is immobilized on a solid substrate or matrix that is insoluble in the fluid in which the other member of the pair resides. The molecular species of the affinity pair bound to the matrix is generally referred to as the ligand, while the liquid soluble

member is generally referred to as the target member. However, it is important to note that these definitions do not impose any restrictions in a strict chemical sense. The vast majority of current ligand immobilization techniques rely on physical or chemical approaches. Physical ligand immobilization involves adsorption or entrapment of the ligand to a suitable support, while the chemical mode of immobilization is characterized by the formation of strong crosslinks or covalent attachments between the ligand and the matrix. It is a requirement that immobilization is accomplished in such a fashion that the capacity of the members of the affinity pair to recognize each other is not adversely affected by the immobilization procedure.

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It is a disadvantage of the currently available physical and chemical techniques for immobilizing ligands that production processes are frequently time consuming and expensive. This is mainly due to the fact that immobilization techniques require the separate production of matrix material and ligands, which in a subsequent step must be coupled. An alternative mode of immobilizing proteins is described in U.S. Patent No. 5,474,925 which documents a biological production system for the immobilization of enzymes in the fibre of cotton plants. This patent discloses what is believed to be the first biologically produced enzyme immobilization system and allows a one step production of matrix and ligand.

Subsequent to immobilization of the ligand on the matrix, a variety of affinity based purification techniques may be employed to accomplish selective binding between the affinity immobilized ligand and the target member. Affinity based purification techniques known in the prior art include perfusion affinity chromatography, affinity repulsion chromatography, hyperdiffusion affinity chromatography, affinity precipitation, membrane affinity partitioning, affinity cross-flow ultrafiltration and affinity precipitation. In the most widely used affinity based purification technique, affinity chromatography, a matrix containing a ligand is coated to, or packed on, the inside of a

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chromatographic column. A complex mixture containing the target member is then applied to the chromatographic column. Ideally, only the target molecules that specifically recognize the ligand bind in a non-covalent fashion to the chromatographic column, while all other molecular species present in the sample pass through the column.

In affinity partitioning, two solutions of substantially different densities are employed. The complex solution containing the target member is mixed with a solution of a different density containing the affinity ligand. Subsequent to mixing, the solutions are left to settle in order to permit the formation of two separate phases. Molecules tend to partition differentially between phases depending on their size, charge and specific interactions with the phase-forming solutions. Ligand-bound target protein selectively partitions to the phase containing the affinity ligand. For example, Coughlin and Baclaski in *Biotechnology Progress*, 1990 6: 307-309 reported the use of the biotin containing organic solution isooctane to transfer avidin from an aqueous solution to the isooctane solution. However, so far applications of affinity partitioning have been limited mainly due to the current lack of availability of suitable affinity matrix substances which can be employed in specific partitioning in two phase systems.

An important factor for the commercial development of biotechnology is the purification of bioproducts, which typically accounts for 50% or more of the total costs (Labrou, N. and Clonis, Y. D. in the *Journal of Biotechnology* 36: 95-119 (1994)). Many protein purification steps rely on column type separation procedures. In particular, large scale high-separation techniques such as column chromatography or batch-type based protein purification techniques are costly. In addition, crude material is less suitable for either column chromatography or batch separations, as contaminants may foul up sedimented resins and plug columns. Thus, affinity matrices are often only employed in a later stage of purification processes where substantial purity is critical, where the proteins are present in extremely dilute concentrations, or where high

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value proteins are required, for example in diagnostic and therapeutic proteins. These and other topics related to the use of affinity technology in biotechnological processes have been reviewed by Labrou, N. and Clonis, Y. D. in the *Journal of Biotechnology* 36: 95-119 (1994).

There is a need in the art to develop novel and economical methods for separating and purifying biological products from complex mixtures. The present inventors have found that subcellular oil storage structures, known as oil bodies, and their associated proteins are useful in this regard.

10 SUMMARY OF THE INVENTION

The present invention relates to a novel versatile biological system for the production of affinity matrices. The present inventors have found that oil bodies and their associated proteins can be used as affinity matrices for the separation of a wide variety of target molecules such as proteins, carbohydrates, lipids, organic molecules, nucleic acids, metals, cells and cell fractions from a sample.

In accordance with the invention, there is provided a method for the separation of a target molecule from a sample comprising: 1) contacting (i) oil bodies that can associate, either directly or indirectly, with the target molecule with (ii) a sample containing the target molecule; and 2) separating the oil bodies associated with the target molecule from the sample. The oil bodies and the sample containing the target molecule are brought into contact in a manner sufficient to allow the oil bodies to associate with the target. Preferably, oil bodies are mixed with the target. If desired, the target molecule may be further separated from the oil bodies.

In one aspect, the target molecule has affinity for, or binds directly to, the oil bodies or oil body protein. Examples of such targets include antibodies or other proteins that bind to oil bodies.

In another aspect, a ligand molecule may be used to associate the target molecule with the oil bodies.

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In one embodiment, the ligand has natural affinity for the oil bodies or oil body protein. In a specific embodiment, the ligand is an antibody that binds the oil body protein. Such an antibody can be used to separate targets having affinity for the ligand antibody such as anti-IgG antibodies or protein A. A bivalent antibody may also be prepared having binding specificities for both the oil body protein and the target. The antibody against the oil body protein may also be fused to a second ligand having affinity for the target.

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In another embodiment, the ligand is covalently attached to the oil bodies or oil body protein. In one embodiment, the ligand is a protein that is chemically conjugated or produced as a fusion protein with the oil body protein (as described in WO 96/21029). In the latter case, the fusion protein is targeted to and expressed on the oil bodies. In one example, the ligand fused to the oil body protein may be hirudin and can be used to purify thrombin. In another example, the ligand fused to the oil body protein may be metallothionein and can be used to separate cadmium from a sample. In a further example, the ligand fused to the oil body protein may be protein A and can be used to separate immunoglobulins. In yet another example, the ligand fused to the oil body protein may be cellulose binding protein and can be used to separate cellulose from a sample.

In another embodiment, the ligand may be covalently attached to the oil bodies. For example, the ligand may be a small organic molecule such as biotin. Biotinylated oil bodies can be used to separate avidin from a sample.

The present invention also includes modified oil bodies for use as an affinity matrix. Accordingly, the present invention includes a composition comprising oil bodies associated with a molecule, such as a ligand molecule or a target molecule. In one embodiment, the composition comprises oil bodies covalently attached to a ligand molecule, such as biotin.

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The present invention also includes an affinity matrix for use in separating a target molecule from a sample, comprising oil bodies that can associate with the target molecule. The affinity matrix may additionally include a ligand molecule associated with the oil bodies, wherein the ligand molecule is capable of associating with the target molecule.

Other objects, features and advantages of the present invention will become apparent from the following detailed description and attached drawings. It should be understood, however, that the detailed description and associated examples are given by way of illustration only, and various changes and modifications thereto falling within the scope of the invention will become apparent to those skilled in the art. In addition, reference is made herein to various publications, patents and patent applications which are hereby incorporated by reference in their entirety.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1. The nucleotide and deduced amino acid sequence of the 18 KDa oleosin from *Arabidopsis thaliana* as shown in SEQ.ID.NO:1 and SEQ.ID.NO:2.

Figure 2. Sequence of an *Arabidopsis* oleosin-hirudin fusion. Indicated are a portion of the oleosin genomic sequence (from base 1-1620 as reported in van Rooijen *et al* 1992, *Plant Mol. Biol.* 18: 1177-1179), a spacer sequence (base 1621-1635, underlined) and the synthetic DNA sequence encoding the mature hirudin variant-2 isoform (base 1636-1833, italicized) This gene fusion is regulated by the 5' upstream region of the *Arabidopsis* oleosin (bases 1-861) and the nopaline synthase termination sequence (base 1855-2109). The sequence is also shown in SEQ.ID.NO:3 and SEQ.ID.NO:4.

Figure 3. Outline of the steps employed in the construction of pCGOBHIRT, containing the entire oleosin-hirudin construct.

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Figure 4. Schematic diagram illustrating the configuration of the oleosin-hirudin fusion protein on the oil body and the binding of thrombin.

Figure 5. NaCl elution profiles of thrombin from wild type and 4A4 oil body matrices transformed with a construct expressing an oleosin-hirudin fusion.

Figure 6. Purification of a horseradish peroxidase conjugated anti-IgG antibody using an anti-oleosin antibody as a ligand. Schematic diagram illustrating the configuration of the oleosin / anti-oleosin / anti-IgG sandwich complex bound to an oil body.

Figure 7. Illustrates specific binding of anti-IgG antibodies to wild type oil bodies complexed with primary anti-oleosin antibodies as a ligand (left) and binding of anti-IgG antibodies to oil bodies which were not complexed with primary antibodies prior to binding with the secondary antibodies (right).

Figure 8. Sequence of an oleosin metallothionein fusion. Indicated are the coding sequence of a *B. napus* oleosin cDNA (bases 1092-1652, van Rooijen, 1993, *Ph.D. Thesis*, University of Calgary), a spacer sequence (bases 1653-1670, underlined) and the human metallothionein gene *mt-II* (bases 1671-1876, Varshney and Gedamu, 1984, *Gene*, 31: 135-145)). The gene fusion is regulated by an *Arabidopsis* oleosin promoter (bases 1-1072) and ubiquitin termination sequence (bases 1870-2361, ubi3'; Kawalleck *et al.*, 1993, *Plant Mol. Biol.* 21: 673-684). The sequence is also shown in SEQ.ID.NO:6 and SEQ.ID.NO:7.

Figure 9. Outline of the steps employed in the construction of pBIOOM3' containing the entire oleosin-metallothionein construct.

Figure 10. Schematic diagram illustrating the configuration of the oleosin-metallothionein fusion protein on the oil body and binding of cadmium ions.

Figure 11. Illustrates the binding (A) and elution (B) of cadmium to an oil body matrix from wildtype B. carinata seeds and B. carinata seeds transformed with a construct expressing oleosin

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metallothionein gene fusion. Shown is the percentage cadmium bound to the oil body fraction of an oil body fraction harvested from transgenic and untransformed seeds. Bars represent average values of 5 replicate experiments (binding) and 3 replicates (elution).

Figure 12. Illustrates the binding of protein A expressing S. aureus cells to oil bodies treated with varying amounts of anti-oleosin IgGs. Bars represent OD_{600} readings obtained following the procedures as described in Example 5 and using varying amounts of IgGs (0 μ l, 3, μ l, 30

μl, 100 μl of added IgG).

Figure 13. Oligonucleotide primers used to amplify the sequence of the *S. aureus* protein A (The sequence is also shown in SEQ.ID.NO:8; The protein sequence is also shown in SEQ.ID.NO:9). Primer BK266, 5'C *TCC ATG* GAT <u>CAA CGC AAT GGT TTA TC</u> 3' (SEQ.ID.NO:10), a *NcoI* site (italicized) and a sequence identical to a portion of the protein A gene as contained within vector pRITZ2T (Pharmacia) (underlined) are indicated. Primer BK267, 5' GC *AAG CTT* CTA <u>ATT TGT TAT CTG CAG GTC</u> 3' (SEQ.ID.NO:11), a *HindIII* site (italicized), a stop codon (bold) and a sequence complementary to a portion of the protein A gene as contained within pRIT2T (Pharmacia) (underlined) are indicated. The PCR product was digested with *NcoI* and *HindIII* and ligated into pCGNOBPGUSA (Van Rooijen and Moloney, 1995, *Plant Physiol.* 109: 1353-1361) from which the *NcoI*-GUS-HindIII fragment had been removed.

fusion (The sequence is also shown in SEQ.ID.NO:12 and the protein sequence is also shown in SEQ.ID.NO:13 and 14). Indicated are a portion of the oleosin genomic sequence (from base 1 - 1626, as reported in van Rooijen et al., 1992 Plant Mol. Biol. 18: 1177-1179), a spacer sequence encoding a thrombin cleavage site (base 1627 - 1647, underlined) and the DNA sequence encoding protein A (base 1648 - 2437, italicized). Expression is regulated by the Arabidopsis 5' upstream region of the

Arabidopsis oleosin (base 1 - 867) and the nopaline synthase terminator region (base 2437 - 2700).

Figure 15. Schematic diagram illustrating the configuration of the oleosin-protein A fusion protein on the oil body and binding of the immunoglobulin.

Figure 16. A western blot illustrating the binding of HRP-conjugated mouse anti-rabbit antibodies to oil body protein extracts obtained from transgenic *B. napus* lines expressing oleosin-protein A fusion proteins. Shown on a Western blot probed with an HRP-conjugated antibody are oil body protein extracts from transgenic lines, opa 30 (lane 3), opa 31 (lane 4), opa 34 (lane 5), opa 36 (lane 6), opa 47 (lane 7), opa 93 (lane 8), all expressing an oleosin-protein A fusion protein and a control untransformed *B. napus* line (lane 9), as well as lysates of *E. coli* DH5α transformed with pRIT2T expressing protein A (lane 2) and untransformed *E. coli* DH5α (lane 1).

Figure 17 illustrates binding and elution of IgGs to oil bodies isolated from wildtype *B. napus* (bn wt) and a transgenic *B. napus* line, expressing an oleosin protein A fusions. Error bars represent the results from 4 independent experiments.

20 <u>DETAILED DESCRIPTION OF THE INVENTION</u>

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As hereinbefore mentioned, the present invention relates to a novel biological affinity matrix system that employs oil bodies and their associated proteins. The affinity matrix is suitable for the highly-efficient separation of specific targets, including proteins, carbohydrates, lipids, nucleic acids, cells and subcellular organelles, metals and ions, from a sample.

The present invention provides a method for the separation of a target molecule from a sample comprising: 1) contacting (i) oil bodies that can associate either directly or indirectly with the target molecule with (ii) a sample containing the target molecule; and 2) separating the oil bodies associated with the target molecule from the sample. The oil

bodies and the sample containing the target molecule are brought into contact in a manner sufficient to allow the oil bodies to associate with the target. Preferably, the oil bodies are mixed with the target. Indirect association of the oil bodies with the target can be effected using a ligand molecule that can associate with both the oil bodies and the target molecule. The ligand therefore serves to bridge or join the oil bodies with the target molecule may be further separated from the oil bodies and the ligand, if present.

Each of the components of the affinity matrix are discussed in turn below.

<u>Targets</u>

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The term "target" as used herein denotes a desired molecule that one wants to purify, isolate or separate from a sample such as a biological mixture. This technology is amenable for use with virtually any target for which a ligand can be obtained or any target that can directly associate with or bind to an oil body or oil body protein. Possible ligand/target pairs include but are not limited to: protein subunit/subunit associations, antibodies/antigens, receptor protein/signal molecules, nucleic acid binding proteins/nucleic acids; lectins/carbohydrates; lipid binding proteins/lipids; ion binding proteins/ions; and ligands to surface epitopes/cells or subcellular organelles. The target may be obtained from any natural source or may be synthesized chemically. If the target is a macromolecule such as a protein or nucleic acid it may also be produced in recombinant form using any suitable expression system such as bacteria, yeast, plant, insect, mammalian, etc.

Ligands

The term "ligand" used herein denotes a molecule that is capable of associating with both the target molecule and the oil bodies or oil body protein (discussed below). The term "associating with" as used herein includes both covalent and non-covalent binding of the ligand to the oil bodies or the target molecule. For example, the ligand molecule

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may be covalently attached to the oil bodies and non-covalently associate with the target (and vice-versa), or the ligand may non-covalently associate with both the oil bodies and the target molecule. The ligand may be any molecule that can bridge the oil bodies or oil body protein and the target molecule and can include a protein, nucleic acid, carbohydrate or small organic molecule. The ligand may be comprised of two molecules, a first molecule that associates with the oil bodies and a second molecule that associates with the target, wherein the first molecule and the second molecule associate with each other.

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The affinity ligand proteins used for this methodology may be derived from naturally-occurring, known ligand pairs such as those listed above. Alternatively, the ligand may be obtained by screening proteins extracted from cells or organisms, synthesized chemically or produced in libraries comprised of combinatorial peptide sequences, antibodies, or expressed DNA sequences.

In one embodiment, the ligand has natural affinity for the oil bodies or the oil body protein. For example, the ligand may be a protein such as an antibody, that has affinity for the oil body protein. The ligand may also be a molecule other than a protein which has natural affinity for the oil body or oil body protein. Such ligands, capable of binding to the oil bodies or oil body protein, may be associated with a second molecule that can bind the target molecule. For example, the ligand molecule may be an antibody conjugated to avidin and can be used to purify biotin from a sample.

In another embodiment, the ligand is covalently linked to the oil bodies or oil body protein by chemical or recombinant means. Chemical means for preparing fusions or conjugates are known in the art and can be used to prepare a ligand-oil body protein fusion. The method used to conjugate the ligand and oil body must be capable of joining the ligand with the oil body protein without interfering with the ability of the ligand to bind to the target molecule. In one example, the ligand may be a small organic molecule such as biotin that is covalently attached to

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the oil bodies. Biotinylated oil bodies can be used to separate avidin from a sample. The present invention also includes modified oil bodies such as biotinylated oil bodies for use as an affinity matrix. Accordingly, the present invention includes a composition comprising oil bodies attached to a molecule, such as a ligand or a target molecule.

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In a preferred embodiment, the ligand is a protein and can be conjugated to the oil body protein using techniques well known in the art. There are several hundred crosslinkers available that can conjugate two proteins. (See for example "Chemistry of Protein Conjugation and Crosslinking". 1991, Shans Wong, CRC Press, Ann Arbor). crosslinker is generally chosen based on the reactive functional groups available or inserted on the ligand. In addition, if there are no reactive groups a photoactivatible crosslinker can be used. In certain instances, it may be desirable to include a spacer between the ligand and the oil-body Crosslinking agents known to the art include the protein. homobifunctional agents: glutaraldehyde, dimethyladipimidate and Bis(diazobenzidine) and the heterobifunctional agents: Maleimidobenzoyl-N-Hydroxysuccinimide and Sulfo-m-Maleimidobenzoyl-N-Hydroxysuccinimide.

A ligand protein-oil body protein fusion may also be prepared using recombinant DNA techniques. In such a case a DNA sequence encoding the ligand is fused to a DNA sequence encoding the oil body protein, resulting in a chimeric DNA molecule that expresses a ligand-oil body protein fusion protein (discussed in greater detail below). In order to prepare a recombinant fusion protein, the sequence of the DNA encoding the ligand must be known or be obtainable. By obtainable it is meant that a DNA sequence sufficient to encode the protein ligand may be deduced from the known amino acid sequence. It is not necessary that the entire gene sequence of the ligand be used provided that a subsequence encoding the binding domain of the protein ligand is known. Therefore, the ligand can include the complete sequence of, or the binding domain from, the specific ligand protein in question.

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If the DNA sequence of the desired ligand is known, the gene may be synthesized chemically using an oligonucleotide synthesizer. Alternatively, the clone carrying the ligand gene may be obtained from either cDNA or genomic libraries containing the gene by probing with a labelled complementary DNA sequence. The gene may also be specifically amplified from the library using gene-specific oligonucleotide primers and the PCR. If the DNA sequence of the desired ligand is not known, then a partial amino acid sequence may be obtained through N-terminal sequencing of the protein (Matsudaira 1987; *J. Biol. Chem.* 262: 10035-10038). Labelled probes may be synthesized based upon the DNA sequences deduced from this amino acid sequence and used to screen cDNA or genomic libraries as described above. The clone carrying the gene may also be identified from a cDNA expression library by probing either with antibodies raised against the protein ligand, or with the target protein.

Ligands may also be uncovered by probing mixtures of proteins with the target. The target can be immobilized on a support matrix and used to screen proteins extracted from cells and tissues or synthesized chemically. Following binding between the ligand protein and the immobilized target, the matrix is separated from the solution and washed. The protein ligand is subsequently eluted from the matrix and the sequence determined as described above. Alternatively, recombinant protein libraries produced by phage display, such as those comprised of combinatorial peptide sequences (Smith, 1985; Science 228: 1315-1317) or antibody repertoires (Griffiths et al., 1994, EMBO J. 13: 3245-3260, Nissim et al., 1994, EMBO J. 13: 692-698) can be screened with the immobilized target. In this case, binding between the protein ligand and the target would enable separation and recovery of the phage expressing the ligand from the large, complex population of phage encoding nonbinding proteins. A two-hybrid system such as that in yeast (Fields and Sternglanz, 1994; Trends Genet. 10: 286-292) might also be used to identify a ligand from an expressed cDNA library. Here, a gene fusion is

constructed between the sequence encoding the target protein and that of a DNA binding protein. Cells containing this construct are transformed with constructs from a cDNA library where the sequences have been fused to that of a transcriptional activator. Binding between ligands derived from the cDNA library with the target protein allows transcription of a reporter gene to occur. Clones expressing the ligand are then recovered.

To specifically uncover a ligand to oil bodies, a complete or partial oleosin protein may be used as target in any of the above methods. Alternatively, it may be possible to employ intact oil bodies for screening protein extracts, synthetic peptides or phage display libraries. In this case, the oil body would serve both as target and immobilization matrix. Using this approach, a wider variety of ligands may be uncovered; that exhibit affinity not only to oleosins, but to other epitopes present on oil bodies.

Oil bodies and Oil Body Proteins

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Oil bodies are small, spherical, subcellular organelles encapsulating stored triacylglycerides, an energy reserve used by many plants. Although they are found in most plants and in different tissues, they are particularly abundant in the seeds of oilseeds where they range in size from under one micron to a few microns in diameter. Oil bodies are comprised of the triacylglycerides surrounded by a half-unit membrane of phospholipids and embedded with a unique type of protein known as an oil body protein. The term "oil body" or "oil bodies" as used herein includes any or all of the triacylglyceride, phospholipid or protein components present in the complete structure. The term "oil body protein" as used herein means a protein that is naturally present in an oil body. In plants, the predominant oil body proteins are termed "oleosins". Oleosins have been cloned and sequenced from many plant sources including corn, rapeseed, carrot and cotton. The oleosin protein appears to be comprised of three domains; the two ends of the protein, Nand C-termini, are largely hydrophilic and reside on the surface of the oil

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body exposed to the cytosol while the highly hydrophobic central core of the oleosin is firmly anchored within the membrane and triacylglyceride. Oleosins from different species represent a small family of proteins showing considerable amino acid sequence conservation, particularly in the central region of protein. Within an individual species, a small number of different isoforms may exist.

Oil bodies from individual species exhibit a roughly uniform size and density which is dependent in part upon the precise protein/ phospholipid/triacylglyceride composition. As a result, they may be simply and rapidly separated from liquids of different densities in which they are suspended. For example, in aqueous media where the density is greater than that of the oil bodies, they will float under the influence of gravity or applied centrifugal force. In 95% ethanol where the density is less than that of the oil bodies, they will sediment under the same conditions. Oil bodies may also be separated from liquids and other solids present in solutions or suspensions by methods that fractionate on the basis of size. For example, the oil bodies from *B. napus* are minimal, approximately 0.5µm in diameter, and thus may be separated from smaller components using a membrane filter with a pore size less than this diameter.

The oil bodies of the subject invention are preferably obtained from a seed plant and more preferably from the group of plant species comprising: thale cress (Arabidopsis thaliana), rapeseed (Brassica spp.), soybean (Glycine max), sunflower (Helianthus annuus), oil palm (Elaeis guineeis), cottonseed (Gossypium spp.), groundnut (Arachis hypogaea), coconut (Cocus nucifera), castor (Ricinus communis), safflower (Carthamus tinctorius), mustard (Brassica spp. and Sinapis alba), coriander (Coriandrum sativum) linseed/flax (Linum usitatissimum), and maize (Zea mays). Plants are grown and allowed to set seed using agricultural cultivation practises well known to a person skilled in the art. After harvesting the seed and removal of foreign material such as stones or seed hulls, for by example sieving, seeds are

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preferably dried and subsequently processed by mechanical pressing, grinding or crushing. The oil body fraction may be obtained from the crushed seed fraction by capitalization on separation techniques which exploit differences in density between the oil body fraction and the aqueous fraction, such as centrifugation, or using size exclusion-based separation techniques, such as membrane filtration, or a combination of both of these. Typically, seeds are thoroughly ground in five volumes of a cold aqueous buffer. A wide variety of buffer compositions may be employed, provided that they do not contain high concentrations of strong organic solvents such as acetone or diethyl ether, as these solvents may disrupt the oil bodies. The solution density of the grinding buffer may be increased with the addition of 0.4-0.6 M sucrose, in order to facilitate washing as described below. The grinding buffer will also typically contain 0.5 M NaCl to help remove soluble proteins that are not integrally bound to the oil body surface.

Following grinding, the homogenate is centrifuged resulting in a pellet of particulate and insoluble matter, an aqueous phase containing soluble components of the seed, and a surface layer comprised of oil bodies with their associated proteins. The oil body layer is skimmed from the surface and thoroughly resuspended in one volume of fresh grinding buffer. It is important that aggregates of oil bodies are dissociated as thoroughly as possible in order to ensure efficient removal of contaminants in the subsequent washing steps. The resuspended oil body preparation is layered under a floatation solution of lower density (e.g. water, aqueous buffer) and centrifuged, again, separating oil body and aqueous phases. This washing procedure is typically repeated at least three times, after which the oil bodies are deemed to be sufficiently free of contaminating soluble proteins as determined by gel electrophoresis. It is not necessary to remove all of the aqueous phase and to the final preparation water or 50 mM Tris-HCl pH 7.5 may be added and if so desired the pH may be lowered to pH 2 or raised to pH 10. Protocols for isolating oil bodies from oil seeds are available in Murphy, D. J. and

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Cummins I., 1989, Phytochemistry, 28: 2063-2069; and in: Jacks, T. J. et al., 1990, JAOCS, 67: 353-361. A preferred protocol is detailed in example 1 of the present specification.

Oil bodies other than those derived from plants may also be used in the present invention. A system functionally equivalent to plant oil bodies and oleosins has been described in bacteria (Pieper-Fürst et al., 1994, J. Bacteriol. 176: 4328), algae (Rossler, P.G., 1988, J. Physiol. (London), 24: 394-400) and fungi (Ting, J. T. et al., 1997, J. Biol Chem. 272: 3699-3706). Oil bodies from these organisms, as well as those that may be discovered in other living cells by a person skilled in the art, may also be employed according to the subject invention.

Affinity Matrices

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As hereinbefore mentioned, the present invention provides a novel affinity matrix system for the purification of a target molecule from a sample. In one embodiment, the affinity matrix comprises oil bodies that can bind a target molecule in a sample. In such an embodiment, the target molecule may be an antibody that can bind an oil body protein. In another embodiment, the affinity matrix comprises oil bodies or oil body proteins and a ligand that is associated with the oil bodies or oil body proteins and has affinity for a target molecule. In such an embodiment, the ligand may be non-covalently or covalently attached to the oil bodies or oil body protein (as described above).

It is an advantage of the present invention that target substances can be purified or removed from samples through non-covalent association with oil bodies followed by oil body separation. A number of different oil body-ligand configurations are possible. Targets with inherent affinity for a specific ligand proteins such as hirudin to thrombin or heavy metals to metallothionein, may be purified or separated with oil bodies containing that ligand fused to an oleosin. Alternatively, a protein target may also be purified or separated with an oil body affinity matrix by fusing the target to an oil body-specific ligand or to a ligand complimentary to that fused to an oleosin. If desired, a

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protease recognition site or chemical cleavage site may be engineered between the ligand and the target protein to enable proteolytic removal of the ligand from the target protein in the course of purification. A multivalent ligand may also be constructed, such as a bivalent singlechain antibody, in which one domain of the ligand has an affinity for an oil body and the other domain(s) exhibits affinity for the target. In this case, neither the oil body nor the target molecule need to be covalently fused to a ligand. Also, concatamers of ligands may be used to increase the affinity of a matrix for a target, or the sequence of a ligand may be mutated to modulate the affinity for a target when such conditions are Further, mixtures of different ligands may be fused to desirable. recover/remove different types of targets simultaneously. Fusions between different ligands may also be constructed to form bridges between different types of targets or between targets and the oil body affinity matrix. Binding to the affinity matrix may also be achieved by forming bridges between ligand or ligand and target sequences, such as Zn++ ions bridging between polyhistidine sequences.

There are several advantages associated with the use of oil body affinity matrices that make them attractive as purification tools. The flexibility in design that is possible through the different configurations described above, enables a matrix to be constructed to best meet the requirements for a specific target. Also, production of the matrix as part of a natural biological process in seeds is extremely costeffective, since purification and immobilization of the ligand are not necessary. In the case of oleosin-ligand fusions, the ligand is immobilized on the oil body as a result of oleosin targeting within the cell, while oil body-specific ligands will naturally associate with the matrix while present in complex mixtures. Natural immobilization of the ligand on the matrix may also be advantageous in that it eliminates the requirement for chemical cross-linking that may compromise the affinity of the ligand for the target. Finally, oil body affinity matrices offer a unique and attractive purification option particularly for large scale

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operations. The ability to separate the matrix through floatation as a loose suspension enables it to be employed with crude material containing what might otherwise be prohibitive amounts of particulate contaminants. The presence of these contaminants will often foul and block conventional solid matrices applied in columns or batch suspensions limiting their use at early stages in the purification process.

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As mentioned previously, in one embodiment of the invention, ligand protein sequences are genetically fused to the oil body protein. In order to prepare such genetic fusions, a chimeric DNA sequence is prepared that encodes an oil body protein-ligand fusion protein and consists of (a) a DNA sequence encoding a sufficient portion of an oil body protein to provide targeting of the fusion protein to the oil bodies and (b) a DNA sequence encoding a sufficient portion of the ligand protein to provide binding of the target. The inventors have determined that, in general, the N-terminus and the hydrophobic core of an oil body protein are sufficient to provide targeting of the fusion protein to the oil bodies. In particular, for oleosins derived from the plant *Arabidopsis thaliana* amino acids 2 through 123 (as shown in SEQ.ID.NO:1) are sufficient in this regard.

The ligand may be fused to either the N- and/or C-terminal end of the oleosin. It may also be possible to construct an internal fusion between the ligand and oleosin or to fuse the ligand between two oleosin proteins. The chimeric DNA sequence encoding an oil body protein fused to a ligand may be transfected into a suitable vector and used to transform a plant. Two types of vectors are routinely employed. The first type of vector is used for the genetic-engineering and assembly of constructs and typically consists of a backbone such as found in the pUC family of vectors, enabling replication in easily-manipulated and maintained gram negative bacteria such as *E. coli*. The second type of vector typified by the Ti and Ri plasmids, specify DNA transfer functions and are used when it is desired that the constructs be introduced into the

plant and stably integrated into its genome via Agrobacterium-mediated transformation.

A typical construct consists, in the 5' to 3' direction, of a regulatory region complete with a promoter capable of directing expression in plants (preferably seed-specific expression), a protein coding region, and a sequence containing a transcriptional termination signal functional in plants. The sequences comprising the construct may be either natural or synthetic or any combination thereof.

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Both non-seed specific promoters, such as the 35-S CaMV promoter (Rothstein et al., 1987; Gene 53: 153-161) and seed-specific promoters such as the phaseolin promoter (Sengupta-Gopalan et al., 1985; PNAS USA 82: 3320-3324) or the Arabidopsis 18 kDa oleosin (Van Rooijen et al., 1992; Plant Mol. Biol. 18: 1177-1179) promoters may be used. In addition to the promoter, the regulatory region contains a ribosome binding site enabling translation of the transcripts in plants and may also contain one or more enhancer sequences, such as the AMV leader (Jobling and Gehrke 1987; Nature 325: 622-625), to increase the expression of product.

The coding region of the construct will typically be comprised of sequences encoding a ligand fused in frame to an oleosin and ending with a translational termination codon. The sequence for the oleosin may be comprised of any DNA sequence, or part thereof, natural or synthetic, sufficient to encode a protein that can be correctly targeted to, and stably expressed on, an oil body. A detailed description of the characteristics of such a sequence has been reported previously in Moloney, 1993; PCT Patent Appl. WO 93/21320 which is hereby incorporated by reference. The sequence may also include introns. The ligand-encoding region may in turn be comprised of any individual, or combination of, ligand sequences identified as described above. If desired, a protease or chemical recognition site may be engineered between the ligand and the target protein to enable proteolytic removal of the ligand from the target protein in the course of purification.

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The region containing the transcriptional termination signal may comprise any such sequence functional in plants such as the nopaline synthase termination sequence and additionally may include enhancer sequences to increase the expression of product.

The various components of the construct are ligated together using conventional methods, typically into a pUC-based vector. This construct may then be introduced into an *Agrobacterium* vector and subsequently into host plants, using one of the transformation procedures outlined below.

A variety of techniques are available for the introduction of DNA into host cells. For example, the chimeric DNA constructs may be introduced into host cells obtained from dicotyledonous plants, such as tobacco, and oleaginous species, such as B. napus using standard Agrobacterium vectors; by a transformation protocol such as that described by Moloney et al., 1989, (Plant Cell Rep., 8: 238-242) or Hinchee et al., 1988, (Bio/Technol., 6: 915-922); or other techniques known to those skilled in the art. For example, the use of T-DNA for transformation of plant cells has received extensive study and is amply described in EPA Serial No. 120,516; Hoekema et al., 1985, (Chapter V, In: The Binary Plant Vector System Offset-drukkerij Kanters B.V., Alblasserdam); Knauf, et al., 1983, (Genetic Analysis of Host Range Expression by Agrobacterium, p. 245, In Molecular Genetics of the Bacteria-Plant Interaction, Puhler, A. ed., Springer-Verlag, NY); and An et al., 1985, (EMBO J., 4: 277-284). Conveniently, explants may be cultivated with A. tumefaciens or A. rhizogenes to allow for transfer of the transcription construct to the plant cells. Following transformation using Agrobacterium the plant cells are dispersed in an appropriate medium for selection, subsequently callus, shoots and eventually plantlets are recovered. The Agrobacterium host will harbour a plasmid comprising the vir genes necessary for transfer of the T-DNA to the plant cells. For injection and electroporation, (see below) disarmed Ti-plasmids (lacking the tumour genes, particularly the T-DNA region) may be introduced into the plant cell.

The use of non-Agrobacterium techniques permits the use of the constructs described herein to obtain transformation and expression in a wide variety of monocotyledonous and dicotyledonous plants and other organisms. These techniques are especially useful for species that are intractable in an Agrobacterium transformation system. Other techniques for gene transfer include biolistics (Sanford, 1988, Trends in Biotech., 6: 299-302), electroporation (Fromm et al., 1985, Proc. Natl. Acad. Sci. USA, 82: 5824-5828; Riggs and Bates, 1986, Proc. Natl. Acad. Sci. USA 83: 5602-5606) or PEG-mediated DNA uptake (Potrykus et al., 1985, Mol. Gen. Genet., 199: 169-177).

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In a specific application, such as to *B. napus*, the host cells targeted to receive recombinant DNA constructs typically will be derived from cotyledonary petioles as described by Moloney *et al.*, (1989, *Plant Cell Rep.*, 8: 238-242). Other examples using commercial oil seeds include cotyledon transformation in soybean explants (Hinchee *et al.*, 1988. *Bio/Technology*, 6: 915-922) and stem transformation of cotton (Umbeck *et al.*, 1981, *Bio/Technology*, 5: 263-266).

Following transformation, the cells, for example as leaf discs, are grown in selective medium. Once shoots begin to emerge, they are excised and placed onto rooting medium. After sufficient roots have formed, the plants are transferred to soil. Putative transformed plants are then tested for presence of a marker. Southern blotting is performed on genomic DNA using an appropriate probe, for example an *A. thaliana* oleosin gene, to show that integration of the desired sequences into the host cell genome has occurred.

The expression cassette will normally be joined to a marker for selection in plant cells. Conveniently, the marker may be resistance to a herbicide, e.g. phosphinothricin or glyphosate, or more particularly an antibiotic, such as kanamycin, G418, bleomycin, hygromycin, chloramphenicol, or the like. The particular marker employed will be one which will allow for selection of transformed cells compared with cells lacking the introduced recombinant DNA.

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The fusion peptide in the expression cassette constructed as described above, expresses at least preferentially in developing seeds. Accordingly, transformed plants grown in accordance with conventional ways, are allowed to set seed. See, for example, McCormick et al. (1986, *Plant Cell Reports*, 5: 81-84). Northern blotting can be carried out using an appropriate gene probe with RNA isolated from tissue in which transcription is expected to occur, such as a seed embryo. The size of the transcripts can then be compared with the predicted size for the fusion protein transcript.

Oil body proteins are then isolated from the seed and analyses performed to determine that the fusion peptide has been expressed. Analyses can be for example by SDS-PAGE. The fusion peptide can be detected using an antibody to the oleosin portion of the fusion peptide. The size of the fusion peptide obtained can then be compared with predicted size of the fusion protein.

Two or more generations of transgenic plants may be grown and either crossed or selfed to allow identification of plants and strains with desired phenotypic characteristics including production of recombinant proteins. It may be desirable to ensure homozygosity of the plants, strains or lines producing recombinant proteins to assure continued inheritance of the recombinant trait. Methods of selecting homozygous plants are well know to those skilled in the art of plant breeding and include recurrent selfing and selection and anther and microspore culture. Homozygous plants may also be obtained by transformation of haploid cells or tissues followed by regeneration of haploid plantlets subsequently converted to diploid plants by any number of known means, (e.g.: treatment with colchicine or other microtubule disrupting agents).

Method of Separating Target Molecules Using the Affinity Matrices

As hereinbefore mentioned, the present invention relates to a method of separating a target molecule from a sample using the above described oil body proteins and in some cases, ligands. In the method of WO 98/27115 PCT/CA97/00951 - 24 -

the invention, oil bodies are mixed with a sample containing the desired target and the interaction between the ligand and target results in the non-covalent association of the target with the oil body. Following centrifugation, the oil bodies and affinity-bound target are separated from the aqueous phase, effectively purifying the target from any contaminants present in the original sample. Repeating the washing step ensures that any remaining contaminants are removed.

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Following their attachment to oil bodies, targets may be eluted under conditions determined empirically for each individual ligand-target pair. Treatment of the bound matrix with the appropriate eluent and centrifugation enables recovery of the purified target in the aqueous phase. If the target is a ligand-protein fusion containing a protease recognition site, then it may be treated with the appropriate protease to remove the ligand. The free ligand may then be separated from the target protein by re-application of the oil body affinity matrix or through conventional protein purification methods.

The chemical and physical properties of the affinity matrix may be varied in at least two ways. Firstly, different plant species contain oil bodies with different oil compositions. For example, coconut is rich in lauric oils (C12), while erucic acid oils (C22) are abundantly present in some Brassica spp. Furthermore, proteins associated with the oil bodies will vary between species. Secondly, the relative amounts of oils may be modified within a particular plant species by applying breeding and genetic engineering techniques or a combination of these known to the skilled artisan. These techniques aim at altering the relative activities of enzymes controlling the metabolic pathways involved in oil synthesis. Through the application of these techniques, seeds with a sophisticated set of different oils are obtainable. For example, breeding efforts have resulted in the development of a rapeseed with a low erucic acid content (Canola) (Bestor, T. H., 1994, Dev. Genet. 15: 458) and plant lines with oils with alterations in the position and number of double bonds, variation in fatty acid chain length and the introduction of desirable functional

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groups have all been generated through genetic engineering (Töpfer et al., 1995, Science, 268: 681-685). Using similar approaches a person skilled in the art will be able to further expand on the presently available sources of oil bodies. Variant oil compositions will result in variant physical and chemical properties of the oil body fraction. Thus by selecting oilseeds or mixtures thereof from different species or plant lines as a source for oil bodies, a broad repertoire of oil body matrices with different textures and viscosities may be acquired.

Applications of Oil Body Affinity Matrices

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Given that it is possible to engineer oil body affinity matrices for several classes of proteins, multiple uses for oil body based affinity matrices are envisioned. Bacteria, fungi, plants and animals all contain proteins which are able to specifically interact with agents such as ions, metals, nucleic acids, sugars, lipids and other proteins. These agents may be immobilized using oil body technology.

The oil body protein affinity matrices can be used to isolate any target molecule that can bind to the oil body protein, either directly or indirectly through a ligand molecule. Examples of target molecules that may be isolated from a sample using the methodology of the present invention include proteins, peptides, organic molecules, lipids, carbohydrates, nucleic acids, cells, cell fragments, viruses and metals. In particular, the inventors have shown that the affinity matrix of the present invention can be used to separate therapeutic proteins (such as thrombin), antibodies, metals (such as cadmium), carbohydrates (such as cellulose), organic molecules (such as biotin) and cells (such as bacterial cells).

Oil body affinity matrices may also be used to separate cells of industrial or medical interest from a mixed population of cells. For example haematopoietic stem cells, which are a subpopulation of blood cells and are used in bone marrow transplantations and in stem cell gene therapies, may be separated from other blood cells using oil body based affinity technology. In recombinant DNA technology it is often required

that cells in which recombinant DNA has been successfully introduced, known as transformed cells, are distinguished and separated from cells which failed to acquire recombinant DNA. Provided that part of the recombinant DNA expresses a cell surface protein which is complementary to a oil body based affinity ligand, it is possible to utilize oil bodies to separate transformed cells from untransformed cells. Oil body affinity technology may also be used to separate cellular organelles such as chloroplasts and mitochondria from other cellular material. Viral particles may also be separated from complex mixtures.

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It is also possible to immobilize a class of proteins known as metalloproteins, which contain prosthetic groups that specifically bind ions. Examples of metalloproteins are haemoglobin, which binds iron, parvalbumbin which binds calcium and metallothionein a protein which binds zinc and other metal ions. It is envisioned that oil bodies could be used to scavenge metals from streams of flowing material, which might be water contaminated with the waste of metals from laboratories and industrial processes. Example 4 given below further illustrates this application. Other examples where proteins may be bioimmobilized and employed in a bioremediation strategy include the removal of phosphates, nitrates and phenols from waste streams. In part this approach may overcome the real or perceived limitations of bacterial bioremediation. In certain instances it may not be practical or necessary to rely on affinity partitioning technology to separate the oil body matrix from the target compound. In these instances, it is envisioned that oil bodies may be immobilized on a solid inert surface which could be a flat surface or the surface of a column. A solution containing the affinity ligand may then be passed over the surface coated with immobilized oil bodies whereupon selective affinity binding occurs. It is envisioned that immobilized oil bodies may be used in pipes and in ponds to assist in bioremediation.

The following examples illustrate various systems in which oil bodies can be used as affinity matrices. It is understood that the

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examples given below are intended to be illustrative rather than limiting.

EXAMPLES

EXAMPLE 1

5 Purification of Thrombin

The following example demonstrates the utility of an oil body affinity matrix for the purification of thrombin. Thrombin is a serine protease which plays a central role in blood coagulation. It cleaves fibrinogen to produce fibrin monomers which polymerize to form the basis of a blood clot (Fenton 1981; Ann. N.Y. Acad. Sci. 370: 468-495). Alfa-thrombin consists of two polypeptide chains of 36 (A-chain) and 259 (B-chain) residues linked by a disulphide bridge. Degen et al. 1983; Biochemistry 22: 2087-2097). Hirudin, which is found in the salivary glands of the medicinal leech Hirudo medicinalis, is a very specific and potent inhibitor of thrombin. This inhibition is a result of the non-covalent binding of hirudin to specific parts of the alfa-thrombin chain. (Stone and Hofsteenge 1986; Biochemistry 25: 4622-4628).

The immobilized ligand is comprised of an isoform of hirudin fused to the 18 kDa *Arabidopsis* oleosin (oil body protein) (Van Rooijen *et al.*, 1992; *Plant Mol. Biol.* 18: 1177-1179). Expression of the construct is regulated by the Arabidopsis 18 kDa oleosin promoter (Van Rooijen *et al.*, 1994; *Plant Mol. Biol.* 18: 1177-1179). The sequence of the oleosin-hirudin fusion is shown in Figure 2 and in SEQ.ID.NO:3.

Oleosin-Hirudin Construct

Oligonucleotide primers were designed based upon the reported sequence for a Brassica napus oleosin gene (Murphy et al. 1991, Biochim. Biophys. Acta 1088: 86-94) and used to amplify a fragment from B. napus genomic DNA through PCR. Using this fragment as a probe, a clone carrying a 15 kbp insert was identified and isolated from a EMBL3 Arabidopsis genomic library. Oligonucleotide primers were used to amplify a fragment from this insert containing the entire oleosin coding sequence and intron together with 840 basepairs of the 5' upstream

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region. The primers were designed so as to eliminate the translational stop codon and to introduce a *PstI* restriction endonuclease recognition site at the 5' end and a *SalI* followed by a *PvuI* site at the 3' end of the fragment. The fragment was end-filled and ligated into the *SmaI* site of the plasmid vector pUC19. A *SalI* - *EcoRI* fragment from plasmid pBI121 (Clontech) comprising the nopaline synthetase terminator sequence was then inserted to generate pOBILT.

A synthetic hirudin variant 2 (HV2) sequence was synthesized based upon reported sequence information (Harvey et al. 1986, Proc. Natl. Acad. Sci. USA 83: 1084-1088) but employing B. napus and Arabidopsis codon usage. The sequence was amplified using four overlapping oligonucleotide primers designed such that the resulting fragment possessed PvuI and SalI sites at the 5' and 3' ends respectively. This fragment was ligated into the SmaI site of the pUC19 plasmid vector to generate pHIR. The PvuI - SalI fragment from pHIR was then inserted into pUCOBILT between the oleosin and terminator sequences to form an in-frame fusion with the oleosin coding region giving pUCOBHIRT. The entire construct was subcloned into pBluescript KS+ (pBIOBHIRT) and then into the PstI site of pCGN1559 plasmid (McBride and Summerfelt, 1990, Plant Mol. Biol. 14: 269-276) carrying a neomycin phosphotransferase gene under control of the 35-S CaMV promoter (pCGOBHIRT). This plasmid was introduced into Agrobacterium tumefaciens. The preparation of this plasmid is shown in Figure 3.

Transformation and Regeneration

Procedures for the transformation of Agrobacterium and plants have been described previously. Agrobacterium tumefaciens was transformed with the above construct through electroporation (Dower et al., 1988; Nucl. Acids Res. 16: 6127-6145). The transformed bacteria were then used to transform cotyledonary explants of Brassica napus, followed by plant regeneration according to the methods of Moloney et al. (1989; Plant Cell Reports 8: 238-242). Transgenic plant were initially identified using a neomycin phosphotransferase assay and subsequently

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confirmed by expression of the oleosin-hirudin fusion as determined through northern and immunoblot analysis.

Preparation of Oil Bodies

Seed from either control (non-transgenic) plants or transgenic plants expressing the oleosin-hirudin fusion were homogenized in five volumes of cold grinding buffer (50 mM Tris-HCl, pH 7.5, 0.4 M sucrose and 0.5 M NaCl) using a polytron operating at high-speed. The homogenate was centrifuged at approximately 10 x g for 30 min. to remove particulate matter and to separate oil bodies from the aqueous phase containing the bulk of soluble seed protein. Oil bodies were skimmed from the surface of the supernatant with a metal spatula and placed in one volume of fresh grinding buffer. To achieve efficient washing in subsequent steps, it was important to ensure that the oil bodies were thoroughly redispersed. This was accomplished by gently re-homogenising the oil bodies in grinding buffer with the polytron operating at low-speed. Using a syringe, the resuspended oil bodies were carefully layered underneath five volumes of cold 50 mM Tris-HCl, pH 7.5 and centrifuged as above. Following centrifugation, the oil bodies were again removed and the washing procedure repeated three times to remove residual contaminating soluble seed proteins. The final washed oil body preparation was resuspended in one volume of cold 50 mM Tris-HCl pH 7.5, redispersed with the polytron, and was then ready for use as an affinity matrix.

Affinity Purification of Thrombin

The purification of thrombin using the oleosin-hirudin fusion protein is shown schematically in Figure 4. In order to evaluate the binding of thrombin, affinity matrices were prepared from transgenic Brassica napus seeds expressing the oleosin-hirudin fusion protein (4A4 seeds) (Parmenter et al. Plant Molecular Biology (1995) 29: 1167-1180) and from wild type Brassica napus cv Westar seeds. Binding of thrombin to both matrices was evaluated. Procedures for the preparation of washed oil bodies from seeds were the same as those described above.

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Solutions containing a range of thrombin activities between 0 and 1 units were mixed with 10 µl of a fixed amount of affinity matrix (prepared from a total of 10 mg of dried seeds; corresponding to approximately 100 μg of total oil body protein) in 500 μl binding buffer (50 mM Tris-HCl (pH 7.5); 0.1% (w/v) BSA). The oil body suspension was then incubated for 30 minutes on ice and centrifuged at 14,000 rpm for 15 minutes at 4°C. The buffer under the oil bodies (termed 'unternatant') containing the unbound, free thrombin was recovered using an hypodermic needle and assayed for thrombin activity as follows. A total of 250 μ l of unternatant was added to 700 µl binding buffer and prewarmed to 37°C. Following addition the of 50 μl of 1 mMthrombin N-p-tosyl-gly-pro-arg-p-nitroanilide (Sigma) to the unternatant, the change in optical density at 405 nanometers was monitored spectrophotometrically for 3 minutes. The concentration of thrombin in the assay mixture was determined employing a standard curve which was constructed using a set of thrombin samples containing known concentrations of thrombin. The values obtained from these assays were used to calculate the concentration bound thrombin assuming:

[bound thrombin] = [total thrombin] - [free thrombin]

The ratio of the concentration of bound over the concentration of free thrombin was plotted as a function of the concentration of bound thrombin (Scatchard plot). From these plots the dissociation constants of the affinity matrix were calculated following standard procedures (Scatchard, G. Ann. N.Y. Acad. Sci. (1949) 57: 660-672) and assuming: $K_a = 1/K_d$. The dissociation constants of the affinity matrices were $3.22 \times 10^{-7} \text{m}$ for wild type and $2.60 \times 10^{-8} \text{m}$ for 4A4 oil bodies.

In order to evaluate the recovery of bound thrombin from the matrices a NaCl gradient was employed. The elution profile of thrombin bound to oleosin-hirudin oil body matrices was compared with the profile from thrombin bound to wildtype oil body matrices.

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Procedures for preparation of wild type oil bodies from wild type Brassica napus cv Westar seeds and for the preparation of oleosin-hirudin oil bodies from Brassica napus 4A4 seeds (Parmenter et al. Plant Molecular Biology (1995) 29: 1167-1180) were identical to those described above. Procedures for binding of thrombin to the matrices were as described above, except 100 µl aliquots of oil bodies were used to bind 0.5 units of thrombin. Oil body suspensions were left on ice for 30 minutes prior to centrifugation for 15 minutes at 4°C and 14,000 rpm. The unternatant was assayed for (unbound) thrombin activity. The oil body matrix was then resuspended in binding buffer to which NaCl was added to a final concentration of 0.05 M. Starting with the 30 minutes incubation of the oil body suspension on ice, the procedure was repeated five times increasing the NaCl concentration in a stepwise fashion. The final NaCl concentrations used were 0.05 M, 0.1 M, 0.2 M, 0.3 M, 0.4 M and 0.6 M. The NaCl concentrations in the thrombin assay were kept constant at 150 mM. Figure 5 shows the elution profiles obtained when wildtype oil bodies and 4A4 oil bodies were used.

EXAMPLE 2

Use of Antibodies as Bivalent Ligands

Antibodies may be used as bivalent ligands by virtue of their affinity both for specific epitopes and for other antibodies or proteins (for example the *Staphylococcus aureus* protein A) which have affinity for immunoglobulins (IgGs). In this example, polyclonal anti-oleosin antibodies serve as a bivalent ligand and antibodies raised in rabbits against the anti-oleosin antibodies serve as the target. This example is illustrated schematically in Figure 6.

Oil bodies were prepared from 5 g of wild type *Brassica* napus cv Westar seeds following the procedure described in Example 1. Subsequently, oil bodies were washed twice with 100 mM glycine (pH 2.5), neutralized through two washes in binding buffer (50 mM Tris-HCl, pH 7.5) and resuspended in 5 ml of binding buffer. A 150 µl aliquot of the

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washed oil body preparation was combined with 500 μl of rabbit serum containing anti-oleosin antibodies (ligand antibodies), diluted 1:10 with binding buffer. The oil body suspension was mixed thoroughly and incubated for 1 h at 4°C with agitation. Following incubation, unbound ligand antibodies were removed from the oil body suspension through three washes with 1 ml of binding buffer. Oil bodies were then combined with 500 μl of serum diluted 1:500 in binding buffer and containing anti-rabbit IgG antibodies (the target antibodies) conjugated with horseradish peroxidase (HRP) as a detection label (Sigma). suspension was mixed and incubated under conditions identical to those used for the anti-oleosin antibody binding. As a control, target antibodies were incubated with oil bodies which had not been previously bound to ligand antibodies. Both samples were subsequently washed four times with 1 ml of binding buffer to remove unbound antibodies. Using binding buffer, the samples were equalized with respect to concentration of oil bodies as determined by measuring sample turbidity spectrophotometrically at 600 nm. To assay for bound target antibody, samples containing 5 µl of oil bodies were mixed with 1 ml of the HRP colorimetric substrate tetramethylbenzidine in 0.01% hydrogen peroxide and reacted for 10 minutes at room temperature. Reactions were stopped by the addition of 500 μl of 1 M $H_2 SO_4$ and the absorbance at 450 nm was determined. Corrections for the presence of residual, unbound target antibody remaining after washing were made by assaying 5 μl of the final wash fraction. The results obtained for control and ligand bound oil body preparations are set forth in Figure 7.

EXAMPLE 3

Use of Oleosin-Specific Ligands

The use of an oleosin-specific ligand represents an alternative to the use of an antibody or genetically-engineered oleosin fusion proteins for the purification of recombinant target proteins. In this case, the target protein is fused to the oleosin-specific ligand and the

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endogenous oleosins present on the oil bodies of non-transgenic seeds serve as the complementary ligand-affinity matrix. In addition to eliminating the requirement for a transgenic line expressing an oleosin fusion, this approach increases the overall capacity of the affinity matrix, since all of the endogenous oleosins may now participate in binding.

Oleosin-specific ligands may be identified and isolated from a peptide phage display library screened with oleosin protein. Since the extreme hydrophobicity of the oleosin central domain can result in aggregation and precipitation of the protein when removed from oil bodies, a mutant protein lacking this domain may be used for screening. This has little effect on the efficacy of the ligand, as only the hydrophillic portions of the oleosin are exposed to the cytoplasm (i.e. the N- and C-termini). Hence, these are the only regions available for binding to a ligand. Once isolated, the ligand may be fused to a common reporter protein, green fluorescent protein (GFP) (Prasher, 1995, *Trends Genet*. 11:320-323), to demonstrate purification.

Removal of the Oleosin Central Domain

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Oligonucleotide primers specific for the *Arabidopsis* oleosin gene described above can be used to amplify an oleosin gene from a *B. napus* cDNA library (van Rooijen 1993, *Ph.D. Thesis*, University of Calgary). Primers flanking sequences encoding the N-terminal 62 amino acids and the C-terminal 55 amino acids, may be used to amplify sequences for the respective N- and C-terminal oleosin domains in separate reactions. Additionally, the primer for the 5' end of the N-terminal domain contains a sequence for a thrombin recognition site to enable cleavage of the fusion protein as described below. The resulting fragment was ligated into the *Smal* site of the bacterial expression vector pEZZ 18 (Pharmacia). This vector contains sequences encoding a signal peptide for protein secretion into the periplasm, and synthetic IgG binding domains derived from protein A to facilitate protein purification, downstream of the multiple cloning site.

Expression and Purification of the Oleosin Deletion Construct

The vector carrying the deletion mutant construct is introduced into E. coli using standard methods and transformants selected. A culture of the transformed bacteria can be induced to express the synthetic protein A-mutant oleosin fusion protein by addition of 1 mM IPTG. Induced cells may be pelleted and resuspended in 5 mM MgSO₄ causing lysis of the periplasmic membrane through osmotic shock. The lysed cells are centrifuged and the supernatant containing the secreted protein is loaded on to a column containing IgG-coupled sepharose. After washing to remove unbound protein, the column is loaded with a buffer containing 50 mM Tris-HCl, pH 7.5, 150 mM NaCl and 1.0U/ml of purified Bovine thrombin (Sigma) to cleave the mutant oleosin from the synthetic protein A. Following incubation at 37°C for 4h, the column is drained and the eluate passed through a column of heparin-coupled sepharose to remove thrombin. The eluate from this column, containing the mutant oleosin protein, is recovered and purity of the protein examined through gel electrophoresis followed by staining with Coomassie blue R250.

Generation of a Peptide Combinatorial Library

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A random peptide combinatorial library may be generated according to the methods of Scott and Smith (1990; Science 249: 386-390). Briefly, the PCR is used to amplify a synthetic DNA fragment containing the degenerate sequence (NNK)₆; where 'N' represents an equal mixture of deoxynucleotides G, A, T, and C, and K represents an equal mixture of deoxynucleotides G and T. The degenerate sequence encodes for hexameric peptides among which are represented every possible combination of the 20 amino acids and amber stop codon. The PCR product is ligated into the gene III sequence of the filamentous bactcophage fUSE and the resulting phagemid introduced into E. coli through electroporation.

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Identification and Isolation of Oleosin-Specific Ligands

The peptide phage display libraries are amplified, concentrated and stored in aliquots of 1012 tdu/ml. Purified mutant oleosin protein is biotinylated using a thiol-cleavable linker (S-S biotin, Pierce) and purified by size exclusion chromatography. Aliquots of the peptide phage display library containing 5x1011 tdu in two ml are screened with the biotinylated protein at a concentration of 50 nM. Phage binding the mutant oleosin protein are recovered using streptavidin-coated paramagnetic beads. Following washing, the phage are eluted through the addition of 50mM dithiothreitol which cleaves the disulphide bond. The eluted phage are then incubated with an excess of log-phase F+ E. coli. Aliquots of the infected cells are plated to determine the phage titre and the remaining cells used in successive rounds of amplification and screening. Following enrichment of the eluted phage by 3-4 orders of magnitude, individual phage are selected and tested for binding to mutant oleosin by direct ELISA. Binding by phage is detected using anti-phage antibodies (Crosby and Schorr, 1995, In Annual Review of Cell Biology). Single stranded DNA is isolated from phage exhibiting binding and the peptide-encoding sequence determined.

20 Affinity Purification with Oleosin-Specific Ligands

The sequence for an oleosin ligand isolated as described above is fused in-frame upstream the sequence for *gfp10* (Prasher *et al.*, 1992, *Gene* 111: 229-233) encoding GFP and the construct ligated into the bacterial expression vector pKK233 (Pharmacia). Soluble protein is extracted through sonication of cells induced to express the ligand-GFP fusion, and adjusted to a concentration of 10 mg/ml in 50 mM Tris-HCl, pH 7.5.

Twenty ml of the protein solution is mixed with 2ml of oil bodies prepared as described above, from seeds of non-transgenic plants. The mixture is incubated at 4°C for 30 min with agitation to allow binding and then centrifuged to separate the oil bodies and soluble fraction. The amount of GFP remaining in the soluble fraction after

removal of oil bodies is determined by fluorescence spectrofluorometry at a wavelength of 508 nm and compared with that in the original bacterial extract. The amount of bound GFP is calculated to determine the capacity of the matrix.

The oil bodies are washed twice in 20 ml of 50 mM Tris-HCl, pH 7.5, resuspended in 2 ml of the same buffer and divided into 20 aliquots of 100 μ l. Conditions for the elution of ligand-GFP fusion protein are determined by adding 1ml of solutions ranging in pH from 2-10 and in NaCl concentration from 0-1 M to different aliquots. After mixing and incubation at 4°C for 30 min, the oil bodies are removed and the soluble fractions collected. The amount of ligand-GFP fusion protein in the soluble fraction is determined by fluorescence spectrophotometry.

EXAMPLE 4

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Removal of Heavy Metal Ions

The following example demonstrates the utility of oil body affinity matrices for the recovery/removal of non-protein targets from complex solutions. For the purpose of this example the metallothionein/Cd++ ligand pair was used. However other metal binding proteins such as phytochelatins (Rauser, 1990; Ann. Rev. Biochem; 59: 61-86) and metal ions including Cu++ and Zn++ could also be used.

Oleosin-Metallothionein Fusion

An oleosin gene from a *B. napus* cDNA library (van Rooijen 1993, *Ph.D. Thesis*, University of Calgary) was amplified through PCR with oligonucleotide primers designed so as to create *Not1* and *Nco1* sites at the 5' and 3' ends of the gene respectively. The resulting fragment was digested and placed into the *Not1/Nco1* sites of pGN to yield plasmid poleGN. The human metallothionein gene, *mt-II* (Varshney and Gedamu, 1984, *Gene*, 31: 135-145) was amplified using oligonucleotide primers designed to create a unique *Not1* site at the 3'-end of the gene. The resulting PCR product was subcloned into the

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blunt-end *EcoRV* site of pBluescript KS+ to form pBSMTC. The *mt-II* gene was then excised from this plasmid and subcloned into the *NcoI/KpnI* sites of poleGN replacing the GUS-NOS region to generate pOLEMTC. The 773 base oleosin-MT fusion of pOLEMTC was excised with *NotI* digestion and inserted into the unique *NotI* site of polePN3' between the oleosin promoter (oleP; Van Rooijen *et al.*, 1992, *Plant Mol. Biol.* 18: 1177-1179) and the *P. crispum ubi4*-2 gene terminator (ubi3'; Kawalleck *et al.*, 1993, *Plant Mol. Biol.* 21: 673-684.) to generate pOOM3'. After the fusion was determined to be in the correct orientation, pOOM3' was digested with *KpnI* to release the oleP-oleMT-ubi3' insert. This expression cassette was inserted at the *KpnI* site of the binary vector pCGN1559 to yield the final construct pBIOOM3'. The sequence of the oleosin-metallothionein fusion is shown in Figure 8 and SEQ.ID.NO.6. The construction of plasmid pB100M3' is shown in Figure 9.

15 Transformation and Regeneration

Transgenic *B. carinata* plants expressing the oleosin-metallothionein fusion were created using transformation and regeneration protocols as described in Example 1.

Oil Body Preparation

Washed oil bodies were prepared from *B. carinata* seeds of transgenic and control plants as described in Example 1.

Removal of Cd++ From Solution Using an Oil Body Affinity Matrix

The use of the oleosin-metallothionein fusion to bind cadmium ions in solution is shown schematically in Figure 10.

A solution of 10 μ M CdCl₂ in 10 mM Tris-HCl, pH 7.2 containing 0.01 μ Ci/ml ¹⁰⁹Cd was prepared. A 1 ml aliquot of this CdCl₂ solution was thoroughly mixed with 100 μ l of washed oil bodies (1.6 mg oil body protein) prepared from seeds expressing the oleosin-metallothionein fusion protein and incubated at 22°C for 1 hr. Following centrifugation for 5′ at 10,000 xg to separate the oil bodies from the aqueous phase and 2 washes in 1 ml of 10 mM Tris-Cl, pH 7.2, the

amount of ¹⁰⁹Cd++ remaining bound to oil body fraction was determined using a gamma-counter (Cobra auto-gamma, Canberra Packard, Canada). An identical experiment was performed with oil bodies from non-transgenic seeds to detect and correct for non-specific binding of Cd ions to the matrix.

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Cd++ ions were eluted from the oil body metallothionein affinity matrix by mixing of the oil body fraction with 1 ml of 100 mM glycine (pH = 3.0) buffer (Pazirandeh et al., 1995; Appl. Microbiol. Biotechn. 43: 1112-1117). Following centrifugation for 5 min. At 10,000 xg, the oil body fraction was removed and assayed for bound Cd++ ions as above. Figure 11 shows Cd binding and elution from the affinity matrix.

EXAMPLE 5

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Separation of Whole Cells

The following example illustrates the capacity of oil bodies to immobilize whole cells. One potential for the use of bacterial cell separation lies in the utility for diagnostics. It is also desirable to separate unique eukaryotic cells such as lymphocytes and stem cells from complex mixtures of cells where the cell type of interest is present in relatively low numbers.

20 Binding of Staphylococcus aureus to oil bodies via protein A

For the purpose of this example, *S. aureus* cells, which express protein A as a surface antigen were mixed with oil bodies with varying amounts of polyclonal anti-oleosin antibodies.

Preparation of oil bodies

Seeds of *B. napus* cv Westar were surface sterilized in bleach, rinsed and ground with a mortar and pestle in grinding buffer (50 mM Tris pH 7.5, 0.4 M sucrose and 100 mM glycine). The homogenate was filtered through Miracloth into sterile 15 ml Corex tubes. The filtered homogenate was then centrifuged for at 4°C for 10 min at 10,000 xg. The oil body fraction was removed and resuspended in 50 mM Tris pH 7.5 and 0.4 M sucrose and washed two times using the same buffer. Aliquots of 1 ml oil bodies were transferred to 1.5 ml Eppendorf tubes

and centrifuged at room temperature for $10 \, \text{min}$ at $16,000 \, \text{xg}$. The oil bodies were washed in $50 \, \text{mM}$ Tris pH $7.5 \, \text{and} \, 0.4 \, \text{M}$ sucrose 5-6 more times until no visible pellet was observed.

Binding of S. aureus cells to anti-oleosin coated oil bodies

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Formalin fixed *S. aureus* cells (Sigma, P-7155) were washed 3-4 times in 50 mM Tris-Cl pH 7.5. and resuspended. Washed oil bodies (300 μ l) and *S. aureus cells* (were mixed with varying amounts of antioleosin IgGs (50 μ l). After mixing and incubating at room temperature for 2 hrs, the mixtures were centrifuged at room temp at 16,000 xg for 5 min. The oil body fraction and unternatant were carefully removed and the cell pellet was washed twice in 1 ml 50 mM Tris-Cl pH 7.5. The walls of the tube were wiped with a tissue to remove traces of oil. Subsequently the drained cell pellets were resuspended in 1 ml of water and the OD₆₀₀ were determined. Figure 12 is a representative experiment showing the decrease in the amount of cells present in the cell pellet as the concentration of anti-IgGs present in the oil-body *S. aureus* mixture increases.

Differential Binding of Two Strains of Staphylococcus aureus.

In this experiment an oil body affinity matrix is employed to demonstrate differential binding of two strains of Staphylococcus aureus. Formalin fixed S. aureus strains, one expressing the IgG binding surface antigen protein A and one lacking protein A, are commercially available from Sigma. Dilute aliquots of both S. aureus strains of equal OD_{550} could be prepared. To each of these aliquots control oil bodies from untransformed plants or oil bodies mixed with anti-oleosin antibodies could be added. Following incubation for an appropriate length of time at an appropriate temperature, the samples could be centrifuged to pellet unbound bacterial cells and to separate the oil body fraction. The oil bodies could be decanted, vortexed and the OD_{550} could be determined. The pellets could be resuspended and the OD_{550} of the unternatant could be determined. It is anticipated that only in the sample containing the S.

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aureus strain expressing protein A and the oil body complexed with anti-oleosin antibodies, fractionation of these cells to the oil body fraction will be observed. Binding of the cells to the oil body could be further demonstrated by lowering of the pH of the oil body fraction. Subsequent to centrifugation the release of cells from the oil bodies could be evidenced by the presence of a pellet and/or an increase in OD_{550} upon resuspension of the pellet.

Separation of Staphylococcus aureus from E. coli

A viable *S. aureus* strain could be mixed with varying quantities of cells of an *E. coli* strain having a specific antibiotic resistance. The mixed bacterial sample could be vortexed with control antibodies and with oil bodies which have been complexed with anti-oleosin antibodies. After incubation for an appropriate length of time and at an appropriate temperature oil bodies could be washed and the unternatant and oil bodies could be directly titrated and selectively plated on blood agar for *S. aureus* growth and on LB plates for *E. coli* growth. The enrichment or actual separation obtained could be determine by an estimate of colony forming units.

Identification of Pathogens Present in Low Concentrations in a Complex Mixture

For diagnostic purposes it is often desirable to concentrate bacterial or viral pathogens which invade human or animal tissues in low numbers. An oil body affinity matrix could be used to enrich for these pathogens, so that they could subsequently be identified and characterized.

Pathogens often specifically bind to human or animal cells through the interaction with a receptor or surface protein. Oleosin could be fused to the human or animal protein ligand and recombinant oil bodies could be employed to immobilize the pathogens. Examples of the formation of protein complexes formed between proteins of human and pathogenic origins known to the prior art include: human fibrinogen or fibrin specific domains which bind to *S. aureus* protein clumping factor

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A (clf-A) (McDevitt et al. 1995; Mol. Microbiol. 16; 895-907); human decay accelerating factor (DAF) to which urinary and intestinal tract pathogenic E. Coli bind (Nowicki et al. 1993: J. Of Experim. Med. 178: 2115-2121); a human cell ligand which is expressed in the carcinoma cell line Caco-2 and which binds uniquely to the 28 kD Klebsiella pneumoniae fimbria protein KPF-28 (Di Maretino et al., 1996; Infect. and Immun. 64: 2263-2266) and human cell extracellular matrix fibronectin specific domains which complex specifically with Streptococcus pyrogenes adhesin (protein F) (Ozeri et al., 1996; EMBO J. 15: 989-998).

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Separation of Small Organic Molecules

This example describes how an oil body affinity matrix may be used for the recovery/removal of small organic molecules from solution. By way of example, the small organic molecule, biotin, is purified using avidin as a ligand.

Construction of Avidin Ligands

Avidin is a protein synthesized by avian species and exhibits an extremely high affinity for biotin, a natural co-factor for many carboxylases. Preparations of purified avidin (commercially available from Sigma) can be conjugated chemically to anti-oleosin antibodies using standard procedures known to those skilled in the art. This approach would yield a bivalent avidin ligand suitable to demonstrate affinity based removal of biotin. Alternatively, an oleosin-avidin gene fusion may be utilized. The gene encoding avidin in chicken (*Gallus gallus*) has been identified and its sequence has been determined (Beattie et al., 1987, Nucl Acids Res. 15: 3595-3606). Based on the sequence the gene for avidin could be synthesized chemically or through the PCR and fused to the B. napus oleosin (van Rooijen, 1993, Ph.D. Thesis, University of Calgary) as described in example 4. Streptavidin, an analogous bacterial biotin binding protein, could also be employed.

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Oil Body Preparation

Washed oil bodies would be prepared from seeds of transgenic plants and/or control plants as described in example 1.

Binding of Bivalent Avidin-Oleosin Ligand

Binding of anti-oleosin antibodies and removal of unbound ligand will be as detailed in example 3.

Removal of Biotin from Solution

Solutions containing known concentrations of biotin could be combined with a fixed amount of oil bodies complexed with an anti-oleosin antibodies conjugated with avidin. Following binding, the mixture would be centrifuged to separate oil body and aqueous fraction. The amount of biotin remaining in the aqueous fraction is determined by competitive ELISA using anti-biotin antibodies conjugated to horse radish peroxidase (HRP). The amount of bound biotin may be calculated assuming:

[bound biotin] = [total biotin] - [free biotin]

From the obtained values, the dissociation constants can be determined as described in example 2. As a control, an identical experiment could be performed with oil bodies bound to anti-oleosin antibodies which have not been conjugated with avidin. If desired, biotin could be released from the oil body-avidin matrix through competitive elution using an excess of 2-(4'-hydroxybenzene) benzoic acid (HABA). Elution may also be aided by employing a genetically engineered mutant of avidin which exhibits a lower affinity for biotin. Such mutants have been described for the analogous biotin binding protein from bacteria, streptavidin (Chilkoti et al., 1995; Bio/Technol. 13: 1198-1204).

EXAMPLE 7

Separation of Carbohydrates

The following example describes the utility of oil body matrices for the recovery of carbohydrates from complex biological

mixtures. In this example the inventors demonstrate that an oil body immobilized cellulase is capable of binding cellulose.

Oleosin-Cellulose Binding Domain Fusion

Several of the cellulases produced by the bacterium Cellulomonas fimi contain discrete cellulose binding domains (CBDs). 5 These CBDs independently bind to cellulose even when they are separated by proteolytic cleavage or genetic manipulation from the catalytic domain of the enzyme. Plasmid pUC18-CBDPT contains a fragment coding for the CBD of the beta-1,4-glucanase (Gilkes et al., 1992, Journal of Biol. Chem. 267: 6743-6749) and could be used to construct an 10 oleosin-CBD gene fusion. A DNA fragment encoding the CBD domain could be isolated from pUC18-CBDPT using appropriate restriction enzymes or using the PCR. Alternatively, the CBDs of other cellulases from C. Fimi or cellulases from other sources could be used. An oleosin gene from B. Napus isolated from a cDNA library (van Rooijen, 1993, 15 Ph.D. Thesis, University of Calgary) was cloned in pGN using the PCR and yielding plasmid pOLEGN as described in example 4. An in-frame gene fusion between the oleosin gene and the CBD gene could be generated using standard molecular techniques known to those skilled in 20 The final construct would comprise the CBD domain translationally fused immediately downstream of the oleosin.

Transformation and Regeneration

In order to introduce the fusion gene construct in plants, it would be subcloned in a binary vector, such as pCGN1559. Transgenic plants which express the oleosin-CBD fusion could be generated as described in example 1.

Oil Body Preparation

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Washed oil bodies could be prepared from the seeds of transgenic and control wild type plants as described in example 1.

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Removal of Cellulose from Solution Using an Oil Body Affinity Matrix

In order to evaluate binding of cellulose to the oil body affinity matrix, the binding capacities of oil bodies of wild type and transgenic plants are compared. Oil bodies could be mixed with appropriately buffered solutions containing a range of cellulose concentrations. The oil body suspension could then be incubated for an appropriate length of time and at an appropriate temperature. Upon centrifugation, the unternatant could be recovered and assayed for cellulose concentrations. The concentrations bound cellulose and free cellulose could be calculated assuming:

[bound cellulose] = [total cellulose] - [free cellulose]

The ratio of the concentration bound over the concentration free cellulose could be plotted as a function of the concentration of bound cellulose. From these plots dissociation constants could be calculated following standard procedures (Scatchard, G. Ann. N. Y. Acad. Sci. (1949) 57: 660-672) and as detailed in example 2.

EXAMPLE 8

Separation of Nucleic Acids

The following example describes a method in which oil bodies are employed to bind single stranded (SS) nucleic acids.

Isolation of Single Stranded Nucleic Acids

A method for capturing SS nucleic acids may be used in diagnostics, such as plant viral disease, or in research applications where non-reannealed SS nucleic acids need to be selectively removed from solutions such as in hybridization reactions for differential screening of expressed genes. Oleosins could be fused with SS DNA or RNA binding proteins or specific domains thereof and could be used to trap SS nucleic acids for identification or further amplification. Well characterized SS nucleic acid binding proteins include: Agrobacterial Ti plasmid Vir E2 protein (Zupan et al., 1995, Plant Physiol. 107: 1041-1047); Tobacco Mosaic Virus (TMV) movement protein P30 (Citovsky et al., 1990; Cell

60: 637-647; Waigmann et al., 1994 Proc Natl. Acad. Sci (USA) 91: 1433-1437); Cauliflower Mosaic Virus coat protein (Thompson et al., 1993; J. Gen. Virol 74: 1141-1148) and E. Coli RecA and single stranded binding (SSB) proteins (Radding, 1991 J. Biol. Chem. 266: 5355-5358).

5 EXAMPLE 9

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Separation of Recombinant Proteins

The following example further demonstrates the utility of an oil body affinity matrix for the purification of recombinant target proteins. For the purpose of this example, the IgG/protein A ligand pair has been chosen. The construct employed consists of a protein A domain which was fused to the 18 kDa *Arabidopsis* oleosin (Van Rooijen *et al.*, 1992; *Plant Mol. Biol.* 18: 1177-1179). Oil bodies containing oleosin-protein A fusion proteins were isolated and used to demonstrate specific binding of rabbit-anti-mouse IgGs conjugated to Horse Raddish Peroxidase (HRP). The configuration of the oleosin-protein A fusion on the oil body and binding of IgG to the fusion is shown in Figure 15.

The Oleosin-Protein A Fusion

A synthetic protein A sequence encoding a protein capable of binding to IgG was synthesized based on reported sequence information (pRIT2T, protein A gene fusion vector; Pharmacia) and was 20 amplified through the PCR. Each primer used in the PCR contained restriction sites 5' to the protein A-specific sequence in order to facilitate cloning. The reverse primer (i.e. the primer in the antisense direction) also contained a translational stop codon following the coding sequence. Fig 13 shows the position of the PCR primers relative to the protein A 25 sequence. (The protein A sequence and the primer sequences are also separately shown in SEQ.ID.NO:8, SEQ.ID.NO:10 and SEQ.ID.NO:11 respectively). The resulting fragment was ligated into a pUC19 plasmid carrying the Arabidopsis oleosin gene comprised of an 867 bp upstream promoter region followed by the coding region (with its associated 30 intron) from which the translational stop codon had been removed. The 3' end of the construct contains the nopaline synthase transcriptional

terminator. A spacer sequence encoding a recognition sequence for the endoprotease thrombin was incorporated immediately downstream of the oleosin coding sequence. The protein A gene sequence was introduced between this spacer sequence and the terminator sequence. In the final expression construct the oleosin and protein A coding regions were fused in the same reading frame. The entire construct (Figure 14 and SEQ.ID.NO:12) was then excised from the pUC19 plasmid and subcloned into the plant transformation vector pCGN1559 (McBride and Summerfelt, 1990, Plant Mol. Biol. 14: 269-276) carrying a neomycin phosphotransferase gene under the control of the 35S CaMV promoter. The resulting plasmid was introduced in Agrobacterium (strain EHA101).

Transformation and Regeneration

Plants were transformed and regenerated as described in example 1. Transgenic plants were initially identified using a neomycin phosphotransferase assay and subsequently confirmed by expression of protein A fusions through immunoblot analysis.

Preparation of Oil Bodies

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Oil bodies from the transgenic *B. napus* and *B. carinata* lines expressing the oleosin-protein A fusion were prepared following the procedure described in example 1.

Binding of Oleosin-Protein A Fusions to IgG

Oil body protein extracts (20 µg/ aliquot) from various transgenic *B. napus* lines expressing oleosin-protein A fusion proteins were subjected to polyacrylamide gelelectrophoresis and subsequently transferred to a PVDF membrane following standard procedures. The membrane was then probed with a HRP-conjugated mouse anti-rabbit antibody and visualised following the procedure as outlined in Antibodies, a laboratory manual (Harlow and Lane, 1988, Cold Spring Harbor). In Figure 16 the stained PVDF membrane is shown. A 50 kDa protein (predicted molecular mass of the oleosin-protein A fusion protein: 48,801 Da) was specifically detected in the protein extracts of all of

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the six transgenic *B. napus* lines tested. Untransformed control plants did not exhibit HRP activity, while the a 30 kDa protein (predicted molecular mass 29,652 Da) was present in a bacterial lysate transformed with pRIT2T encoding protein A and undetectable in the untransformed lysate.

5 Binding and Elution of IgGs to Oil Bodies Containing Oleosin-Protein A Fusion Proteins

Washed oil bodies (10 mg/ml protein) were prepared from wildtype B. napus and a transgenic B. napus line transformed with a construct expressing an oleosin-protein A fusion protein as described in example 1 and suspended in 10 mM Tris-Cl pH 8.0. A volume of 2 μ l (\pm 34 μ g) of HRP-conjugated rabbit anti-mouse antibodies (Sigma, cat no A9044) was added to 500 μ l of the washed oil body preparation and the suspension was incubated for 1 hr at room temperature or overnight at 4°C. The samples were then centrifuged for 15 min at 16,000 \times g and the undernatant was removed. Subsequently, the oil bodies were thoroughly resuspended in 500 μ l 10 mM Tris-Cl pH 8.0 using a pestle. This washing step in Tris-Cl was repeated 4 times (henceforth termed washed oil body preparation). A 5 μ l aliquot from the washed oil body preparation was washed a fifth time and then assayed for HRP activity.

HRP assays were carried out by adding 1 μ l of the washed oil body preparation to 1 ml of HRP assay mix (9.8 ml of 0.1 M NaOAc, 0.2 ml of 2.5 mg/ml Trimethylbenzidine in DMSO, 4 μ l H₂O₂) and incubating the mixture for 5 min at room temperature. The reaction was then stopped by adding 0.5 ml 1M H₂SO₄. The samples were filtered through a 0.22 μ m filter and subsequently the OD₄₅₀'s were determined spectrophotometrically.

In order to elute the IgGs from the oil bodies, the washed oil body preparation was resuspended in 100 mM glycine pH 3.0 and centrifuged for 15 min at 16,000 xg and incubated for 30' at room temperature. Following neutralization in 500 μ l 100 mM Tris-Cl pH 8.0,

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both the oil body fraction and the eluate were assayed for HRP activity as above. The binding and elution of IgGs to oil bodies from weld type *B. napus* and transgenic *B. napus* expressing an oleosin protein A fusion, are illustrated in Figure 17.

All publications, patents and patent applications are herein incorporated by reference in their entirety to the same extent as if each individual publication, patent or patent application was specifically and individually indicated to be incorporated by reference in its entirety.

SEQUENCE LISTING

(1) GENERAL INFORMATION:

- (i) APPLICANTS:
 - (A) NAME: SemBioSys Genetics Inc.
 - (B) STREET: 609-14 Street, N.W.
 - (C) CITY: Calgary
 (D) STATE: Alberta

 - (E) COUNTRY: Canada
 - (F) POSTAL CODE: T2N 2A1
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 - (H) TELEFAX NO.: (403) 220-0704
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 (D) STATE: Alberta
 - (E) COUNTRY: Canada
 - (F) POSTAL CODE: T3A 5N5

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 (B) STREET: #302, 332 6th Avenue N.W.
 - (C) CITY: Calgary
 - (D) STATE: Alberta
 - (E) COUNTRY: Canada
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 - (B) STREET: 3223 Bearspaw Drive N.W.
 - (C) CITY: Calgary
 - (D) STATE: Alberta
 - (E) COUNTRY: Canada
 - (F) POSTAL CODE: T2L 1T1
- (ii) TITLE OF INVENTION: Oil Bodies and Associated Proteins as Affinity Matrices
- (iii) NUMBER OF SEQUENCES: 14
- (iv) CORRESPONDENCE ADDRESS:

 - (A) ADDRESSEE: BERESKIN & PARR
 (B) STREET: 40 King Street West
 - (C) CITY: Toronto
 - (D) STATE: Ontario
 - (E) COUNTRY: Canada (F) ZIP: M5H 3Y2

 - (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk

 - (B) COMPUTER: IBM PC compatible (C) OPERATING SYSTEM: PC-DOS/MS-DOS (D) SOFTWARE: PatentIn Release #1.0, Version #1.30
- (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER:
 - (B) FILING DATE:
 - (C) CLASSIFICATION:
- (vii) PRIOR APPLICATION DATA:
 - (A) APPLICATION NUMBER: US 08/767,026
 - (B) FILING DATE: 16-DEC-1997
- (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: Gravelle, Micheline

- 50 -

(B) REGISTRATION NUMBER: 40,261 (C) REFERENCE/DOCKET NUMBER: 9369-050

(ix) TELECOMMUNICATION INFORMATION:

- (A) TELEPHONE: (416) 364-7311
- (B) TELEFAX: (416) 361-1398

(2) INFORMATION FOR SEQ ID NO:1:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 522 base pairs

 - (B) TYPE: nucleic acid (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Oleosin From Arabidopsis Thaliana
- (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 1..522

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

ATG Met 1	GCG Ala	GAT Asp	ACA Thr	GCT Ala 5	AGA Arg	GGA Gly	ACC Thr	CAT His	CAC His 10	GAT Asp	ATC Ile	ATC Ile	GGC Gly	AGA Arg 15	GAC Asp		48
CAG Gln	TAC Tyr	CCG Pro	ATG Met 20	ATG Met	GGC Gly	CGA Arg	GAC Asp	CGA Arg 25	GAC Asp	CAG Gln	TAC Tyr	CAG Gln	ATG Met 30	TCC Ser	GGA Gly		96
CGA Arg	GGA Gly	TCT Ser 35	GAC Asp	TAC Tyr	TCC Ser	AAG Lys	TCT Ser 40	AGG Arg	CAG Gln	ATT Ile	GCT Ala	AAA Lys 45	GCT Ala	GCA Ala	ACT Thr		144
GCT Ala	GTC Val 50	ACA Thr	GCT Ala	GGT Gly	GGT Gly	TCC Ser 55	CTC Leu	CTT Leu	GTT Val	CTC Leu	TCC Ser 60	AGC Ser	CTT Leu	ACC Thr	CTT Leu		192
GTT Val 65	GGA Gly	ACT Thr	GTC Val	ATA Ile	GCT Ala 70	TTG Leu	ACT Thr	GTT Val	GCA Ala	ACA Thr 75	CCT Pro	CTG Leu	CTC Leu	GTT Val	ATC Ile 80		240
TTC Phe	AGC Ser	CCA Pro	ATC	CTT Leu 85	GTC Val	CCG Pro	GCT Ala	CTC Leu	ATC Ile 90	ACA Thr	GTT Val	GCA Ala	CTC Leu	CTC Leu 95	ATC Ile		288
ACC Thr	GGT Gly	TTT Phe	CTT Leu 100	TCC Ser	TCT Ser	GGA Gly	GGG Gly	TTT Phe 105	GGC Gly	ATT Ile	GCC Ala	GCT Ala	ATA Ile 110	ACC Thr	GTT Val		336
TTC Phe	TCT Ser	TGG Trp 115	ATT Ile	TAC Tyr	AAG Lys	TAC Tyr	GCA Ala 120	ACG Thr	GGA Gly	GAG Glu	CAC His	CCA Pro 125	CAG Gln	GGA Gly	TCA Ser	Ü	384
GAC Asp	AAG Lys 130	TTG Leu	GAC Asp	AGT Ser	GCA Ala	AGG Arg 135	ATG Met	AAG Lys	TTG Leu	GGA Gly	AGC Ser 140	AAA Lys	GCT Ala	CAG Gln	GAT Asp		432
CTG Leu	AAA Lys	GAC Asp	AGA Arg	GCT Ala	CAG Gln	TAC Tyr	TAC Tyr	GGA Gly	CAG Gln	CAA Gln	CAT His	ACT Thr	GGT Gly	GGG Gly	GAA Glu		480

145 150 155 160

CAT GAC CGT GAC CGT ACT CGT GGT GGC CAG CAC ACT ACT TAA His Asp Arg Asp Arg Thr Arg Gly Gly Gln His Thr Thr * 165

522

- (2) INFORMATION FOR SEQ ID NO:2:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 174 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Met Ala Asp Thr Ala Arg Gly Thr His His Asp Ile Ile Gly Arg Asp

1 15

Gln Tyr Pro Met Met Gly Arg Asp Arg Asp Gln Tyr Gln Met Ser Gly 20 25 30

Arg Gly Ser Asp Tyr Ser Lys Ser Arg Gln Ile Ala Lys Ala Ala Thr 35 40 45

Ala Val Thr Ala Gly Gly Ser Leu Leu Val Leu Ser Ser Leu Thr Leu 50 60

Val Gly Thr Val Ile Ala Leu Thr Val Ala Thr Pro Leu Leu Val Ile 65 70 75 80

Phe Ser Pro Ile Leu Val Pro Ala Leu Ile Thr Val Ala Leu Leu Ile 85 90 95

Thr Gly Phe Leu Ser Ser Gly Gly Phe Gly Ile Ala Ala Ile Thr Val 100 105 110

Phe Ser Trp Ile Tyr Lys Tyr Ala Thr Gly Glu His Pro Gln Gly Ser 115 120 125

Asp Lys Leu Asp Ser Ala Arg Met Lys Leu Gly Ser Lys Ala Gln Asp 130 140

Leu Lys Asp Arg Ala Gln Tyr Tyr Gly Gln Gln His Thr Gly Gly Glu 145 155 160

His Asp Arg Asp Arg Thr Arg Gly Gly Gln His Thr Thr \star 165 $170 \cdot$

- (2) INFORMATION FOR SEQ ID NO:3:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 2115 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: cDNA
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Oleosin Hirudin Fusion
 - (ix) FEATURE:
 - (A) NAME/KEY: CDS

(B) LOCATION: 862..1215

(ix) FEATURE:

(A) NAME/KEY: CDS (B) LOCATION: 1456..1833

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

	(X	1) S	EQUE	ENCE	DESC	RIPI	: NOI	SEC	OI (NO:3	3:					
CT												TAAC	CTAC	CTC	CACTCC	= 60
CA	GAAA	CAAC	CGG	CGCC	AAA	TTGC	CGGA	T TA	GCTG	ACCI	G AA	GACG	GAAC	ATC	ATCGTC	120
GG'	TCCT	TGGG	CGA	TTGC	GGC	GGAA	GATG	GG I	'CAGC	TTGG	G CI	TGAG	GACG	AGA	.CCCGAAT	120
CG.	AGTC	TGTT	GAA	AGGT	TGT	TCAT	TGGG	AT T	TGTA	TACG	G AG	ATTG	GTCG	TCG	AGAGGTT	180
TG	AGGG.	AAAG	GAC	AAAT	GGG	TTTG	GCTC	TG G	AGAA	AGAG	A GT	GCGG	باستاست	AGA	GAGAGAA	240
TTO	GAGA	GGTT	TAG.	AGAG.	AGA	TGCG	GCGG	CG A	TGAC	GGGA	G GA	GAGA	CCAC	GAG	GACCTGC	300
ATT	PATC	AAAG	CAG'	TGAC	GTG (ĢTGA	\ATT'	TG G.	AACT'	TTTA	A GA	GGCA	COME	CAT	TTATTAT	360
TTC	TAT	CCAT	TTT	CTTC	ATT (GTTC:	ΓAGA	AT G'	TCGC	GGAA(CAA	،سسس ۲۰۰۰	מממד	λ CTI	AAATCCT	420
AAA	TTTT	TOT	AAT	TTTG	TTG (CCAAT	AGTO	GG A	ratg:	rggg	cer	מית ביד. מית ביד	מאמי	CANC	CTATTG	480
AAG	GCC	AAA	CCC	ATAC	rga (CGAGO	CCA	AA GO	GTTCC	3TTTT	י פכנ	_ 	מאאכ	MMM.	GGTTCG	540
ATG	CCAA	CGC	CAC	ATTCT	rga c	CTAC	GCAZ	LA AZ	ACAZ	ACCI	r cord	יחחחת	IAIG	1110	CTCCTCT	600
CGT	TAAC	ACA	TGC	AGCGC	CT (CATO	GTGA	C GC	ייים ב	מסמי	CCT		AAT.	AGAC	GCATGA	660
TGT	CTCC	ATT	GACA	CGTC	AC I	TCTC	GTCT	יר כיז	الماليات	יייא איי	י ארחא	. GGCC	TAC	AATT	CTCCTA	720
CCT	CTTC	CAA	AATA	TATA	CA C	АТСТ	لىششىك 	'G 21	יר א איזי		. ALF	TOTA	ACA	AACA	CTCCTA TTCTCT	780
СТА	GTAA	ACA	AGAA	CAAA	.AA A	АТС	GCG	CAT	י ארא	C1C1	CAT	TCAA	AAT	CTCA	TTCTCT CAC	840
						1	1114	nsp	, 1111	A1a 5	Arg	GLY	Thr	His	His 10	891
			-	15		0111	+ Y L	FIO	20	Mer	GIY	Arg	Asp	Arg 25	GAC Asp	939
CAG Gln	TAC Tyr	CAG Gln	ATG Met 30	TCC Ser	GGA Gly	CGA Arg	GGA Gly	TCT Ser 35	GAC Asp	TAC Tyr	TCC Ser	AAG Lys	TCT Ser 40	AGG Arg	CAĠ Gln	987
ATT Ile	GCT Ala	AAA Lys 45	GCT Ala	GCA Ala	ACT Thr	GCT Ala	GTC Val 50	ACA Thr	GCT Ala	GGT Gly	GGT Gly	TCC Ser 55	CTC Leu	CTT Leu	GTT Val	1035
CTC Leu	TCC Ser 60	AGC Ser	CTT Leu	ACC Thr		GTT Val 65	C T 3	ACT Thr	GTC Val	ATA Ile	GCT Ala 70	TTG Leu	ACT Thr	GTT Val	GCA Ala	1083
75					80	- 110	261	FIO	TIE	85	Val	Pro	GCT Ala	Leu	Ile 90	1131
		÷		95			Oly	riie	100	ser	Ser	Gly	GGG Gly	Phe 105	GGC Gly	1179
ATT	GCC	GCT	ATA	ACC	GTT	TTC	TCT	TGG	ATT	TAC	AAG	TAAG	CACA	.CA		1225

Ile Ala Ala Ile Thr Val Phe Ser Trp Ile Tyr Lys 110 115

TTT	ATCA	TCT	TACT	TCAT	AA Ţ	TTTG	TGCA	А ТА	TGTG	CATG	CAT	GTGT	TGA	GCCA	GTAGCT	,	1285
TTG	GATC	AAT	TTTT'	TTGG	TC G	AATA	ACAA	A TG	TAAC	AATA	AGA	AATT	GCA	AATT	CTAGGG		1345
AAC.	ATTT	GGT	TAAC'	ГААА	TA C	GAAA'	TTTG.	A CC	TAGC	TAGC	TTG	AATG'	TGT	CTGT	GTATAT		1405
CAT	CTAT.	АТА	GGTA	AAAT	GC T	TGGT.	ATGA	T AC	CTAT	TGAT	TGT	GAAT.		TAC Tyr 1			1461
ACG Thr	GGA Gly	GAG Glu 5	CAC His	CCA Pro	CAG Gln	GGA Gly	TCA Ser 10	GAC Asp	AAG Lys	TTG Leu	GAC Asp	AGT Ser 15	GCA Ala	AGG Arg	ATG Met		1509
AAG Lys	TTG Leu 20	GGA Gly	AGC Ser	AAA Lys	GCT Ala	CAG Gln 25	GAT Asp	CTG Leu	AAA Lys	GAC Asp	AGA Arg 30	GCT Ala	CAG Gln	TAC Tyr	TAC Tyr		1557
GGA Gly 35	GIII	CAA Gln	CAT His	ACT Thr	GGT Gly 40	TGG Trp	GAA Glu	CAT His	GAC Asp	CGT Arg 45	GAC Asp	CGT Arg	ACT Thr	CGT Arg	GGT Gly 50		1605
GGC	CAG Gln	CAC His	ACT Thr	ACT Thr 55	GCG Ala	ATC Ile	GAA Glu	GGG Gly	AGA Arg 60	ATC Ile	ACT Thr	TAC Tyr	ACT Thr	GAC Asp 65	TGT Cys		1653
ACT Thr	GAA Glu	TCT Ser	GGA Gly 70	CAG Gln	AAC Asn	CTC Leu	TGT Cys	CTC Leu 75	TGT Cys	GAA Glu	GGA Gly	TCT Ser	AAC Asn 80	GTT Val	TGT Cys		1701
GGA Gly	AAG Lys	GGA Gly 85	AAC Asn	AAG Lys	TGT Cys	ATC Ile	CTC Leu 90	GGA Gly	TCT Ser	AAC Asn	GGA Gly	AAG Lys 95	GGA Gly	AAC Asn	CAG Gln		1749
TGT Cys	GTT Val 100	ACT Thr	GGA Gly	GAA Glu	GGA Gly	ACT Thr 105	CCA Pro	AAC Asn	CCA Pro	GAA Glu	TCT Ser 110	CAC His	AAC Asn	AAC Asn	GGA Gly	:	1797
GAC Asp 115	TTC Phe	GAA Glu	GAA Glu	ATC Ile	CCT Pro 120	GAA Glu	GAA Glu	TAC Tyr	CTC Leu	CAG Gln 125	TAA *	GTCG	ACTO	CTA		:	18.43
GACG	GATO	TC C	CGAT	CGTI	C AA	ACAT	TTGG	CAA	TAAA	GTT	TCTT	'AAGA	TT C	SAATC	CTGTT	1	1903
GCCG	GTCT	TG C	GATG	ATTA	T CA	TATA	ATTI	CTG	TTGA	TTA	ACGT	TAAG	CA I	GTAA	TAATT	1	1963
AACA	TGTA	AT C	CATG	ACGI	TA T	TTAT	GAGA	TGG	GTTT	TTA	TGAT	TAGA	GT C	CCGC	AATTA	2	2023
TACA	ATTT.	AT A	rcece	ATAG	A AA	ACAA	ААТА	TAG	CGCG	CAA	ACTA	.GGAT	AA A	TATT	CGCGC	2	2083
GCGG	TGTC	AT C	TATG	TTAC	T AG	ATCG	GAAT	TC						•		2	2115

(2) INFORMATION FOR SEQ ID NO:4:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 118 amino acids
 (B) TYPE: amino acid
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

Met Ala Asp Thr Ala Arg Gly Thr His His Asp Ile Ile Gly Arg Asp

Gln Tyr Pro Met Met Gly Arg Asp Arg Asp Gln Tyr Gln Met Ser Gly

Arg Gly Ser Asp Tyr Ser Lys Ser Arg Gln Ile Ala Lys Ala Ala Thr

Ala Val Thr Ala Gly Gly Ser Leu Leu Val Leu Ser Ser Leu Thr Leu

Val Gly Thr Val Ile Ala Leu Thr Val Ala Thr Pro Leu Leu Val Ile

Phe Ser Pro Ile Leu Val Pro Ala Leu Ile Thr Val Ala Leu Leu Ile

Thr Gly Phe Leu Ser Ser Gly Gly Phe Gly Ile Ala Ala Ile Thr Val

Phe Ser Trp Ile Tyr Lys

(2) INFORMATION FOR SEQ ID NO:5:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 126 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

Tyr Ala Thr Gly Glu His Pro Gln Gly Ser Asp Lys Leu Asp Ser Ala

Arg Met Lys Leu Gly Ser Lys Ala Gln Asp Leu Lys Asp Arg Ala Gln

Tyr Tyr Gly Gln Gln His Thr Gly Trp Glu His Asp Arg Asp Arg Thr 35 40 45

Arg Gly Gly Gln His Thr Thr Ala Ile Glu Gly Arg Ile Thr Tyr Thr 50 60

Asp Cys Thr Glu Ser Gly Gln Asn Leu Cys Leu Cys Glu Gly Ser Asn 65 70 75 80

Val Cys Gly Lys Gly Asn Lys Cys Ile Leu Gly Ser Asn Gly Lys Gly

Asn Gln Cys Val Thr Gly Glu Gly Thr Pro Asn Pro Glu Ser His Asn 105

Asn Gly Asp Phe Glu Glu Ile Pro Glu Glu Tyr Leu Gln 120

(2) INFORMATION FOR SEQ ID NO:6:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 2366 base pairs

 - (B) TYPE: nucleic acid (C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Oleosin - Metallothionein Fusion

(ix) FEATURE:

(A) NAME/KEY: CDS
(B) LOCATION: 1092..1856

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

DECEMBER 1150. SEQ ID NO. 6:	
GAGCTCAAAT ACGATCTGAT ACTGATAACG TCTAGATTTT TAGGGTTAAA GCAATCAATC	60
ACCTGACGAT TCAAGGTGGT TGGATCATGA CGATTCCAGA AAACATCAAG CAAGCTCTCA	120
AAGCTACACT CTTTGGGATC ATACTGAACT CTAACAACCT CGTTATGTCC CGTAGTGCCA	180
GTACAGACAT CCTCGTAACT CGGATTATGC ACGATGCCAT GGCTATACCC AACCTCGGTC	240
TTGGTCACAC CAGGAACTCT CTGGTAAGCT AGCTCCACTC CCCAGAAACA ACCGGCGCCA	300
AATTGCCGGA ATTGCTGACC TGAAGACGGA ACATCATCGT CGGGTCCTTG GGCGATTGCG	360
GCGGAAGATG GGTCAGCTTG GGCTTGAGGA CGAGACCCGA ATCGAGTCTG TTGAAAGGTT	420
GTTCATTGGG ATTTGTATAC GGAGATTGGT CGTCGAGAGG TTTGAGGGAA AGGACAAATG	480
GGTTTGGCTC TGGAGAAAGA GAGTGCGGCT TTAGAGAGAG AATTGAGAGG TTTAGAGAGA	540
GATGCGGCGG CGATGACGGG AGGAGAGCG ACGAGGACCT GCATTATCAA AGCAGTGACG	600
TGGTGAAATT TGGAACTTTT AAGAGGCAGA TAGATTTATT ATTTGTATCC ATTTTCTTCA	660
TTGTTCTAGA ATGTCGCGGA ACAAATTTTA AAACTAAATC CTAAATTTTT CTAATTTTGT	720
TGCCAATAGT GGATATGTGG GCCGTATAGA AGGAATCTAT TGAAGGCCCA AACCCATACT	780
GACGAGCCCA AAGGTTCGTT TTGCGTTTTA TGTTTCGGTT CGATGCCAAC GCCACATTCT	840
GAGCTAGGCA AAAAACAAAC GTGTCTTTGA ATAGACTCCT CTCGTTAACA CATGCAGCGG	900
CTGCATGGTG ACGCCATTAA CACGTGGCCT ACAATTGCAT GATGTCTCCA TTGACACGTG	960
ACTTCTCGTC TCCTTTCTTA ATATATCTAA CAAACACTCC TACCTCTTCC AAAATATATA	1020
CACATCTTTT TGATCAATCT CTCATTCAAA ATCTCATTCT CTCTAGTAAA CAGGATCCCC	1080
CTCGCGGCCG C ATG GCG GAT ACA GCT AGA ACC CAT CAC GAT GTC ACA AGT Met Ala Asp Thr Ala Arg Thr His His Asp Val Thr Ser 1 5 10	1130
CGA GAT CAG TAT CCC CGA GAC CGA GAC CAG TAT TCT ATG ATC GGT CGA Arg Asp Gln Tyr Pro Arg Asp Asp Gln Tyr Ser Met Ile Gly Arg 15 20 25	1178
GAC CGT GAC CAG TAC TCT ATG ATG GGC CGA GAC CGA GAC CAG TAC AAC Asp Arg Asp Gln Tyr Ser Met Met Gly Arg Asp Arg Asp Gln Tyr Asn 35	1226
ATG TAT GGT CGA GAC TAC TCC AAG TCT AGA CAG ATT GCT AAG GCT GTT Met Tyr Gly Arg Asp Tyr Ser Lys Ser Arg Gln Ile Ala Lys Ala Val 50 55 60	1274

ACC	GCA	GTC	ACG	GCG	GGT	GGG	TCC	· СТС	י ריייים	CTC	CTC	maa			ACC	
			65	****	. Gry		ser	70	Leu	Val	Leu	Ser	Ser 75	Leu	Thr	1322
CTT Leu	GTT Val	GGT Gly 80		GTC Val	ATT Ile	GCT Ala	TTG Leu 85	Thr	GTT Val	GCC Ala	ACT Thr	CCA Pro 90	CTC Leu	CTC Leu	GTT Val	1370
ATC Ile	TTT Phe 95	AGC Ser	CCA Pro	ATC Ile	CTC Leu	GTG Val 100	CCG Pro	GCT Ala	CTC Leu	ATC Ile	ACC Thr 105	GTA Val	GCA Ala	CTT Leu	CTC Leu	1418
ATC Ile 110	ACT Thr	GGC Gly	TTT Phe	CTC Leu	TCC Ser 115	TCT Ser	GGT Gly	GGG Gly	TTT Phe	GCC Ala 120	ATT Ile	GCA Ala	GCT Ala	ATA Ile	ACC Thr 125	1466
GTC Val	TTC Phe	TCC Ser	TGG Trp	ATC Ile 130	TAT Tyr	AAG Lys	TAC Tyr	GCA Ala	ACG Thr 135	GGA Gly	GAG Glu	CAC His	CCA Pro	CAG Gln 140	GGG Gly	1514
TCA Ser	GAT Asp	AAG Lys	TTG Leu 145	GAC Asp	AGT Ser	GCA Ala	AGG Arg	ATG Met 150	AAG Lys	CTG Leu	GGA Gly	ACC Thr	AAA Lys 155	GCT Ala	CAG Gln	1562
GAT Asp	ATT Ile	AAA Lys 160	GAC Asp	AGA Arg	GCT Ala	CAA Gln	TAC Tyr 165	TAC Tyr	GGA Gly	CAG Gln	CAA Gln	CAT His 170	ACA Thr	GGT Gly	GGT Gly	1610
GAG Glu	CAT His 175	GAC Asp	CGT Arg	GAC Asp	CGT Arg	ACT Thr 180	CGT Arg	GGT Gly	GGC Gly	CAG Gln	CAC His 185	ACT Thr	ACT Thr	CTC Leu	GTT Val	1658
CCA Pro 190	CGA Arg	GGA Gly	TCC Ser	ATG Met	GAT Asp 195	CCC Pro	AAC Asn	TGC Cys	TCC Ser	TGT Cys 200	GCC Ala	GCC Ala	AGT Ser	GAC Asp	TCC Ser 205	1706
TGC Cys	ACC Thr	TGC Cys	• • • •	GGC Gly 210	TCC Ser	TGC Cys	AAG Lys	TGC Cys	AAA Lys 215	GAG Glu	TGC Cys	AAA Lys	Cys	ACC Thr 220	TCC Ser	1754
TGC Cys	AAG Lys	-70	AGC Ser 225	TGC Cys	TGC Cys	TCC Ser	Cys	TGT Cys 230	CCT Pro	GTG Val	GGC Gly	Cys .	GCC Ala 235	AAG Lys	TGT Cys	1802
GCC Ala		GGC Gly 240	TGC . Cys	11C	Cys.	AAA (Lys (GIA '	GCG Ala	TCG (Ser /	GAC . Asp	Lys (rgc A Cys S 250	AGC ' Ser (IGC '	IGT Cys	1850
GCC Ala	TGA * 255	GCGG	CCGC	GA G	GGCT	GCAG	A ATO	GAGT	TCCA	AGA'	rggt	rtg 1	rgaco	GAAG'	ΓT	1906
AGTT	GGTT	GT T	TTTA	TGGA	A CT	TTGT:	TAA	GCT'	TGTA.	ATG '	rgga <i>i</i>	\AGA.	AC G	rgtgo	GCTTT	1966
															ГССТА	
															GCCTG	
															TGTAG	
															TAAT	
															AAAT	
CATT	ATAG'	TA GO	CGTC	TTG(G TCT	rgtgi	TCA	TTG	GTTGA	AC A	AAAGG	CACA	C TC	CACTI	GGAG	2326

ATGCCGTCTC CACTGATATT TGAACAAAGA ATTCGGTACC

2366

(2) INFORMATION FOR SEQ ID NO:7:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 255 amino acids(B) TYPE: amino acid

 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

Met Ala Asp Thr Ala Arg Thr His His Asp Val Thr Ser Arg Asp Gln

Tyr Pro Arg Asp Arg Asp Gln Tyr Ser Met Ile Gly Arg Asp Arg Asp

Gln Tyr Ser Met Met Gly Arg Asp Arg Asp Gln Tyr Asn Met Tyr Gly

Arg Asp Tyr Ser Lys Ser Arg Gln Ile Ala Lys Ala Val Thr Ala Val

Thr Ala Gly Gly Ser Leu Leu Val Leu Ser Ser Leu Thr Leu Val Gly

Thr Val Ile Ala Leu Thr Val Ala Thr Pro Leu Leu Val Ile Phe Ser

Pro Ile Leu Val Pro Ala Leu Ile Thr Val Ala Leu Leu Ile Thr Gly

Phe Leu Ser Ser Gly Gly Phe Ala Ile Ala Ile Thr Val Phe Ser

Trp Ile Tyr Lys Tyr Ala Thr Gly Glu His Pro Gln Gly Ser Asp Lys

Leu Asp Ser Ala Arg Met Lys Leu Gly Thr Lys Ala Gln Asp Ile Lys

Asp Arg Ala Gln Tyr Tyr Gly Gln Gln His Thr Gly Gly Glu His Asp

Arg Asp Arg Thr Arg Gly Gly Gln His Thr Thr Leu Val Pro Arg Gly

Ser Met Asp Pro Asn Cys Ser Cys Ala Ala Ser Asp Ser Cys Thr Cys 195 200 205

Ala GÏy Ser Cys Lys Cys Lys Glu Cys Lys Cys Thr Ser Cys Lys Lys 210 220

Ser Cys Cys Ser Cys Cys Pro Val Gly Cys Ala Lys Cys Ala Gln Gly 225 230 235 240

Cys Ile Cys Lys Gly Ala Ser Asp Lys Cys Ser Cys Cys Ala 245

(2) INFORMATION FOR SEQ ID NO:8:

(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 804 base pairs - 58 -

- (B) TYPE: nucleic acid(C) STRANDEDNESS: single(D) TOPOLOGY: linear
- (ii) MOLECULE TYPÉ: cDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Protein A Primers
- (ix) FEATURE:

 - (A) NAME/KEY: CDS (B) LOCATION: 5..796
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

	,,,,,		.gon.													
CTC	C AT Me	G GA t As 1	T CA p Gl	A CC n Ar	C AA g As	T GG n Gl 5	T TT y Ph	T AT e Il	C CA e Gl	n Se	C CI r Le 0	T AA u Ly	A GA 'S As	T GA	T CCA p Pro 15	49
AGC Ser	CAA Gln	AGT Ser	GCT Ala	AAC Asn 20	val	TTA Leu	GGT Gly	GAA Glu	GCT Ala 25	Gln	AAA Lys	CTT Leu	AAT Asn	GAC Asp 30	TCT Ser	97
CAA Gln	GCT Ala	CCA Pro	AAA Lys 35	Ald	GAT Asp	GCG Ala	CAA Gln	CAA Gln 40	AAT Asn	AAC Asn	TTC Phe	AAC Asn	AAA Lys 45	GAT Asp	CAA Gln	145
CAA Gln	AGC Ser	GCC Ala 50	TTC Phe	TAT Tyr	GAA Glu	ATC Ile	TTG Leu 55	AAC Asn	ATG Met	CCT Pro	AAC Asn	TTA Leu 60	AAC Asn	GAA Glu	GCG Ala	193
CAA Gln	CGT Arg 65	AAC Asn	GGC Gly	TTC Phe	ATT Ile	CAA Gln 70	AGT Ser	CTT Leu	AAA Lys	GAC Asp	GAC Asp 75	CCA Pro	AGC Ser	CAA Gln	AGC Ser	241
ACT Thr 80	AAC Asn	GTT Val	TTA Leu	GGT Gly	GAA Glu 85	GCT Ala	AAA Lys	AAA Lys	TTA Leu	AAC Asn 90	GAA Glu	TCT Ser	CAA Gln	GCA Ala	CCG Pro 95	289
AAA Lys	GCT Ala	GAT Asp	AAC Asn	AAT Asn 100	TTC Phe	AAC Asn	AAA Lys	GAA Glu	CAA Gln 105	CAA Gln	AAT Așn	GCT Ala	TTC Phe	TAT Tyr 110	GAA Glu	337
ATC Ile	TTG Leu	AAT Asn	ATG Met 115	CCT Pro	AAC Asn	TTA Leu	AAC Asn	GAA Glu 120	GAA Glu	CAA Gln	CGC Arg	AAT Asn	GGT Gly 125	TTC Phe	ATC Ile	385
CAA Gln	AGC Ser	TTA Leu 130	AAA Lys	GAT Asp	GAC Asp	CCA Pro	AGC Ser 135	CAA Gln	AGT Ser	GCT Ala	AAC Asn	CTA Leu 140	TTG Leu	TCA Ser	GAA Glu	433
GCT Ala	AAA Lys 145	AAG Lys	TTA Leu	AAT Asn	GAA Glu	TCT Ser 150	CAA Gln	GCA Ala	CCG Pro	AAA Lys	GCG Ala 155	GAT Asp	AAC Asn	AAA Lys	TTC Phe	481
AAC Asn 160	AAA Lys	GAA Glu	CAA Gln	CAA Gln	AAT Asn 165	GCT Ala	TTC Phe	TAT Tyr	GAA Glu	ATC Ile 170	TTA Leu	CAT His	TTA Leu	CCT Pro	AAC Asn 175	529
TTA Leu	AAC Asn	GAA Glu	GAA Glu	CAA Gln 180	CGC Arg	AAT Asn	GGT Gly	TTC Phe	ATC Ile 185	CAA Gln	AGC Ser	CTA Leu	AAA Lys	GAT Asp 190	GAC Asp	577
CCA Pro	AGC Ser	CAA Gln	AGC Ser	GCT Ala	AAC Asn	CTT Leu	TTA Leu	GCA Ala	GAA Glu	GCT Ala	AAA Lys	AAG Lys	CTA Leu	AAT Asn	GAT Asp	625

- 59 -

			195					200					205			
GCT Ala	CAA Gln	GCA Ala 210	CCA Pro	AAA Lys	GCT Aļa	GAC Asp	AAC Asn 215	AAA Lys	TTC Phe	AAC Asn	AAA Lys	GAA Glu 220	CAA Gln	CAA G1n	AAT Asn	673
GCT Ala	TTC Phe 225	TAT Tyr	GAA Glu	ATT Ile	TTA Leu	CAT His 230	TTA Leu	CCT Pro	AAC Asn	TTA Leu	ACT Thr 235	GAA Glu	GAA Glu	CAA Gln	CGT Arg	721
AAC Asn 240	GGC Gly	TTC Phe	ATC Ile	CAA Gln	AGC Ser 245	CTT Leu	AAA Lys	GAC Asp	GAT Asp	CCG Pro 250	GGG Gly	AAT Asn	TCC Ser	CGG Arg	GGA Gly 255	769
TCC Ser	GTC Val	GAC Asp	CTG Leu	CAG Gln 260	ATA Ile	ACA Thr	AAT Asn	TAG *	AAGO	TTGC	:					804
(2)	INFO	RMAT	NOI	FOR	SEQ	ID N	10 : 9 :									
	((i) S	EQUE (A)	ENCE LEN	CHAR IGTH :	RACTE 264	RIST ami	'ICS: no a	cids	:						

- (B) TYPE: amino acid
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9: Met Asp Gln Arg Asn Gly Phe Ile Gln Ser Leu Lys Asp Asp Pro Ser Gln Ser Ala Asn Val Leu Gly Glu Ala Gln Lys Leu Asn Asp Ser Gln Ala Pro Lys Ala Asp Ala Gln Gln Asn Asn Phe Asn Lys Asp Gln Gln 35 40 45Ser Ala Phe Tyr Glu Ile Leu Asn Met Pro Asn Leu Asn Glu Ala Gln Arg Asn Gly Phe Ile Gln Ser Leu Lys Asp Asp Pro Ser Gln Ser Thr 65 70 75 80 Asn Val Leu Gly Glu Ala Lys Lys Leu Asn Glu Ser Gln Ala Pro Lys Ala Asp Asn Asn Phe Asn Lys Glu Gln Gln Asn Ala Phe Tyr Glu Ile Leu Asn Met Pro Asn Leu Asn Glu Glu Gln Arg Asn Gly Phe Ile Gln Ser Leu Lys Asp Asp Pro Ser Gln Ser Ala Asn Leu Leu Ser Glu Ala Lys Lys Leu Asn Glu Ser Gln Ala Pro Lys Ala Asp Asn Lys Phe Asn Lys Glu Gln Gln Asn Ala Phe Tyr Glu Ile Leu His Leu Pro Asn Leu

Asn Glu Glu Gln Arg Asn Gly Phe Ile Gln Ser Leu Lys Asp Asp Pro 185

29

- Ser Gln Ser Ala Asn Leu Leu Ala Glu Ala Lys Lys Leu Asn Asp Ala 200
- Gln Ala Pro Lys Ala Asp Asn Lys Phe Asn Lys Glu Gln Gln Asn Ala
- Phe Tyr Glu Ile Leu His Leu Pro Asn Leu Thr Glu Glu Gln Arg Asn 230
- Gly Phe Ile Gln Ser Leu Lys Asp Asp Pro Gly Asn Ser Arg Gly Ser 250

Val Asp Leu Gln Ile Thr Asn *

- (2) INFORMATION FOR SEQ ID NO:10:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 27 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: cDNA
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Primer Bk 266
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:

CTCCATGGAT CAACGCAATG GTTTATC

(2) INFORMATION FOR SEQ ID NO:11:

- (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 29 base pairs

 - (B) TYPE: nucleic acid (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Primer Bk267
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:

GCAAGCTTCT AATTTGTTAT CTGCAGGTC

(2) INFORMATION FOR SEQ ID NO:12:

- (i) SEQUENCE CHARACTERÍSTICS:
 - (A) LENGTH: 2709 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (ix) FEATURE:
 - (A) NAME/KEY: CDS

(ix) FEATURE:

(A) NAME/KEY: CDS
(B) LOCATION: 1462..2436

(B) LOCATION: 868..1220

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:

(XI) SEQ	UENCE	DESCI	RIPTI	ON:	SEQ	ID N	0:12	:					
CCATGGC	TAT A	CCCAAC	CTC C	GTCT	TGGT	C AC	ACCA	.GGAA	CTC	TCTG	GTA	AGCI	AGCTCC	60
ACTCCCC.	AGA A	ACAACO	GGC G	CCAA	ATTG	C CG	GAAT	TGCT	GAC	CTGA	AGA	CGGA	ACATCA	120
TCGTCGG	GTC C	TTGGGC	GAT 1	rgcgg	CGGA	A GA	TGGG	TCAG	CTT	GGGC	TTG	AGGA	CGAGAC	180
CCGAATC	GAG T	CTGTTG	AAA C	GTTG	TTCA	T TG	GGAT	TTGT	ATA	CGGA	.GAT	TGGT	CGTCGA	240
GAGGTTT	GAG G	GAAAGG	ACA A	ATGG	GTTT	G GC	TCTG	GAGA	AAG	AGAG	TGC	GGCT	TTAGAG	300
AGAGAAT'	TGA G	AGGTTT	'AGA G	AGAG	ATGC	G GC	GGCG	ATGA	CGG	GAGG	AGA	GACG	ACGAGG	360
ACCTGCA'	TTA TO	CAAAGO	AGT G	ACGT	GGTG.	A AA	TTTG	GAAC	ттт	TAAG	AGG	CAGA	TAGATT	420
TATTATT	TGT A	TCCATI	TTC I	TCAT	TGTT	C TA	GAAT	GTCG	CGG	AACA	ААТ	TTTA	АААСТА	480
AATCCTA	AAT T	TTTCTA	ATT T	TGTT	GCCA	A TA	GTGG.	ATAT	GTG	GGCC	GTA	TAGA	AGGAAT	540
CTATTGA	AGG C	CCAAAC	CCA T	'ACTG	ACGA	G CC	CAAA	GGTT	CGT	TTTG	CGT	TTTA	TGTTTC	600
GGTTCGA	TGC · C	AACGCC	ACA T	TCTG.	AGCT	A GG	CAAA	AAAC	AAA	CGTG	TCT	TTGA.	ATAGAC	660
TCCTCTCC	GTT A	ACACAT	GCA G	CGGC	TGCA:	r GGʻ	TGAC	GCCA	TTA	ACAC	GTG	GCCT.	ACAATT	720
GCATGATO	GTC TO	CCATTG	ACA C	GTGA	CTTC	r cgʻ	TCTC	CTTT	CTT	AATA'	TAT	CTAA	CAAACA	780
CTCCTAC	CTC T	rccaaa	АТА Т	ATAC.	ACATO	TT'	TTTG	ATCA	ATC'	rctc.	ATT ·	CAAA	ATCTCA	840
TTCTCTCT	TAG TA	AAACAA	GAA C	AAAA	AA A. Me	FG GG et Al	CG GA	AT AC	CA GO	CT AG la A: 5	GA G rg G	GA AG	CC hr	891
CAT CAC His His 10	GAT A	ATC AT Ile Il	C GGC e Gly	AGA Arg 15	GAC Asp	CAG Gln	TAC Tyr	CCG Pro	ATG Met 20	ATG Met	GGC Gly	CGA Arg	GAC Asp	939
CGA GAC Arg Asp 25	CAG 1	TAC CA Tyr Gl	G ATG n Met 30	Ser	GGA Gly	CGA Arg	GGA Gly	TCT Ser 35	GAC Asp	TAC Tyr	TCC Ser	AAG Lys	TCT Ser 40	987
AGG CAG Arg Gln	ATT (GCT AA Ala Ly 4	s Ala	GCA Ala	ACT Thr	GCT Ala	GTC Val 50	ACA Thr	GCT Ala	GGT Gly	GGT Gly	TCC Ser 55	CTC Leu	1035
CTT GTT Leu Val	CTC 1	CC AG Ser Se 60	C CTT r Leu	ACC Thr	CTT Leu	GTT Val 65	GGA Gly	Thr	Val	Ile	GCT Ala 70	Leu	ACT Thr	1083
GTT GCA Val Ala	ACA C Thr F 75	CCT CT Pro Le	G CTC u Leu	GTT Val	ATC Ile 80	TTC Phe	AGC Ser	CCA Pro	ATC Ile	CTT Leu 85	GTC Val	CCG Pro	GCT Ala	1131
CTC ATC Leu Ile	ACA C	GTT GC Val Al	A CTC a Leu	Leu	ATC Ile	ACC Thr	GGT Gly	TTT Phe	Leu	TCC Ser	TCT Ser	GGA Gly	GGG Gly	1179
90				95					100					

Phe Gly Ile Ala Ala Ile Thr Val Phe Ser Trp Ile Tyr GTAAGCACAC ATTTATCATC TTACTTCATA ATTTTGTGCA ATATGTGCAT GCATGTGTTG 1280 AGCCAGTAGC TTTGGATCAA TTTTTTTGGT CGAATAACAA ATGTAACAAT AAGAAATTGC 1340 AAATTCTAGG GAACATTTGG TTAACTAAAT ACGAAATTTG ACCTAGCTAG CTTGAATGTG 1400 TCTGTGTATA TCATCTATAT AGGTAAAATG CTTGGTATGA TACCTATTGA TTGTGAATAG 1460 G TAC GCA ACG GGA GAG CAC CCA CAG GGA TCA GAC AAG TTG GAC AGT Tyr Ala Thr Gly Glu His Pro Gln Gly Ser Asp Lys Leu Asp Ser 1506 GCA AGG ATG AAG TTG GGA AGC AAA GCT CAG GAT CTG AAA GAC AGA GCT Ala Arg Met Lys Leu Gly Ser Lys Ala Gln Asp Leu Lys Asp Arg Ala 1554 CAG TAC TAC GGA CAG CAA CAT ACT GGT GGG GAA CAT GAC CGT GAC CGT Gln Tyr Tyr Gly Gln Gln His Thr Gly Gly Glu His Asp Arg Asp Arg 1602 40 ACT CGT GGT GGC CAG CAC ACT ACT CTC GTT CCA CGA GGA TCC ATG GAT Thr Arg Gly Gly Gln His Thr Thr Leu Val Pro Arg Gly Ser Met Asp 1650 CAA CGC AAT GGT TTT ATC CAA AGC CTT AAA GAT GAT CCA AGC CAA AGT Gln Arg Asn Gly Phe Ile Gln Ser Leu Lys Asp Asp Pro Ser Gln Ser 1698 GCT AAC GTT TTA GGT GAA GCT CAA AAA CTT AAT GAC TCT CAA GCT CCA Ala Asn Val Leu Gly Glu Ala Gln Lys Leu Asn Asp Ser Gln Ala Pro 1746 AAA GCT GAT GCG CAA CAA AAT AAC TTC AAC AAA GAT CAA CAA AGC GCC Lys Ala Asp Ala Gln Gln Asn Asn Phe Asn Lys Asp Gln Gln Ser Ala 1794 105 TTC TAT GAA ATC TTG AAC ATG CCT AAC TTA AAC GAA GCG CAA CGT AAC Phe Tyr Glu Ile Leu Asn Met Pro Asn Leu Asn Glu Ala Gln Arg Asn 1842 GGC TTC ATT CAA AGT CTT AAA GAC GAC CCA AGC CAA AGC ACT AAC GTT Gly Phe Ile Gln Ser Leu Lys Asp Asp Pro Ser Gln Ser Thr Asn Val 1890 135 TTA GGT GAA GCT AAA AAA TTA AAC GAA TCT CAA GCA CCG AAA GCT GAT Leu Gly Glu Ala Lys Lys Leu Asn Glu Ser Gln Ala Pro Lys Ala Asp 1938 150 AAC AAT TTC AAC AAA GAA CAA CAA AAT GCT TTC TAT GAA ATC TTG AAT Asn Asn Phe Asn Lys Glu Gln Gln Asn Ala Phe Tyr Glu Ile Leu Asn 1986 165 170 ATG CCT AAC TTA AAC GAA GAA CAA CGC AAT GGT TTC ATC CAA AGC TTA Met Pro Asn Leu Asn Glu Glu Gln Arg Asn Gly Phe Ile Gln Ser Leu 2034 180 AAA GAT GAC CCA AGC CAA AGT GCT AAC CTA TTG TCA GAA GCT AAA AAG Lys Asp Asp Pro Ser Gln Ser Ala Asn Leu Leu Ser Glu Ala Lys Lys 2082 195 TTA AAT GAA TCT CAA GCA CCG AAA GCG GAT AAC AAA TTC AAC AAA GAA Leu Asn Glu Ser Gln Ala Pro Lys Ala Asp Asn Lys Phe Asn Lys Glu 2130

CAA CAA AAT GCT TTC TAT GAA ATC TTA CAT TTA CCT AAC TTA AAC GAA 2178 Gln Gln Asn Ala Phe Tyr Glu Ile Leu His Leu Pro Asn Leu Asn Glu GAA CAA CGC AAT GGT TTC ATC CAA AGC CTA AAA GAT GAC CCA AGC CAA Glu Gln Arg Asn Gly Phe Ile Gln Ser Leu Lys Asp Asp Pro Ser Gln 2226 245 AGC GCT AAC CTT TTA GCA GAA GCT AAA AAG CTA AAT GAT GCT CAA GCA 2274 Ser Ala Asn Leu Leu Ala Glu Ala Lys Lys Leu Asn Asp Ala Gln Ala CCA AAA GCT GAC AAC AAA TTC AAC AAA GAA CAA CAA AAT GCT TTC TAT Pro Lys Ala Asp Asn Lys Phe Asn Lys Glu Gln Gln Asn Ala Phe Tyr 2322 280 GAA ATT TTA CAT TTA CCT AAC TTA ACT GAA GAA CAA CGT AAC GGC TTC Glu Ile Leu His Leu Pro Asn Leu Thr Glu Glu Gln Arg Asn Gly Phe 2370 295 300 ATC CAA AGC CTT AAA GAC GAT CCG GGG AAT TCC CGG GGA TCC GTC GAC Ile Gln Ser Leu Lys Asp Asp Pro Gly Asn Ser Arg Gly Ser Val Asp 2418 310 315 CTG CAG ATA ACA AAT TAG AAGCTTGCAT GCCTGCAGGT CGATCGTTCA 2466 Leu Gln Ile Thr Asn * AACATTTGGC AATAAAGTTT CTTAAGATTG AATCCTGTTG CCGGTCTTGC GATGATTATC 2526 ATATAATTTC TGTTGAATTA CGTTAAGCAT GTAATAATTA ACATGTAATG CATGACGTTA 2586 TTTATGAGAT GGGTTTTTAT GATTAGAGTC CCGCAATTAT ACATTTAATA CGCGATAGAA 2646 AACAAAATAT AGCGCGCAAA CTAGGATAAA TTATCGCGCG CGGTGTCATC TATGTTACTA 2706 GAT 2709

(2) INFORMATION FOR SEQ ID NO:13:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 117 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:

Met Ala Asp Thr Ala Arg Gly Thr His His Asp Ile Ile Gly Arg Asp
1 5 10 15

Gln Tyr Pro Met Met Gly Arg Asp Arg Asp Gln Tyr Gln Met Ser Gly
20 25 30

Arg Gly Ser Asp Tyr Ser Lys Ser Arg Gln Ile Ala Lys Ala Ala Thr 35 40 45

Ala Val Thr Ala Gly Gly Ser Leu Leu Val Leu Ser Ser Leu Thr Leu 50 55 60

Val Gly Thr Val Ile Ala Leu Thr Val Ala Thr Pro Leu Leu Val Ile 65 70 75 80

Phe Ser Pro Ile Leu Val Pro Ala Leu Ile Thr Val Ala Leu Leu Ile

- 64 -

85 90 95

Thr Gly Phe Leu Ser Ser Gly Gly Phe Gly Ile Ala Ala Ile Thr Val 105

Phe Ser Trp Ile Tyr

(2) INFORMATION FOR SEQ ID NO:14:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 325 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:

Tyr Ala Thr Gly Glu His Pro Gln Gly Ser Asp Lys Leu Asp Ser Ala

Arg Met Lys Leu Gly Ser Lys Ala Gln Asp Leu Lys Asp Arg Ala Gln

Tyr Gly Gln Gln His Thr Gly Gly Glu His Asp Arg Asp Arg Thr 35 40 45

Arg Gly Gly Gln His Thr Thr Leu Val Pro Arg Gly Ser Met Asp Gln

Arg Asn Gly Phe Ile Gln Ser Leu Lys Asp Asp Pro Ser Gln Ser Ala 65 70 75 80

Asn Val Leu Gly Glu Ala Gln Lys Leu Asn Asp Ser Gln Ala Pro Lys

Ala Asp Ala Gln Gln Asn Asn Phe Asn Lys Asp Gln Gln Ser Ala Phe

Tyr Glu Ile Leu Asn Met Pro Asn Leu Asn Glu Ala Gln Arg Asn Gly

Phe Ile Gln Ser Leu Lys Asp Asp Pro Ser Gln Ser Thr Asn Val Leu 140

Gly Glu Ala Lys Lys Leu Asn Glu Ser Gln Ala Pro Lys Ala Asp Asn

Asn Phe Asn Lys Glu Gln Gln Asn Ala Phe Tyr Glu Ile Leu Asn Met

Pro Asn Leu Asn Glu Glu Gln Arg Asn Gly Phe Ile Gln Ser Leu Lys

Asp Asp Pro Ser Gln Ser Ala Asn Leu Leu Ser Glu Ala Lys Lys Leu

Asn Glu Ser Gln Ala Pro Lys Ala Asp Asn Lys Phe Asn Lys Glu Gln

Gln Asn Ala Phe Tyr Glu Ile Leu His Leu Pro Asn Leu Asn Glu Glu

Gln Arg Asn Gly Phe Ile Gln Ser Leu Lys Asp Asp Pro Ser Gln Ser

245 250 255 Ala Asn Leu Leu Ala Glu Ala Lys Lys Leu Asn Asp Ala Gln Ala Pro260 . Lys Ala Asp Asn Lys Phe Asn Lys Glu Gln Gln Asn Ala Phe Tyr Glu Ile Leu His Leu Pro Asn Leu Thr Glu Glu Gln Arg Asn Gly Phe Ile 290 295 300 Gln Ser Leu Lys Asp Asp Pro Gly Asn Ser Arg Gly Ser Val Asp Leu 305 310 315 320 Gln Ile Thr Asn

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WE CLAIM:

- 1. A method for the separation of a target molecule from a sample comprising:
- 1) contacting (i) oil bodies with (ii) a sample containing the target molecule to allow the target molecule to associate with the oil bodies; and
 - 2) separating the oil bodies associated with the target molecule from the sample.
- 2. A method according to claim 1 further comprising a ligand molecule that associates with the oil bodies and the target molecule.
 - 3. A method according to claim 2 wherein the ligand molecule is covalently attached to the oil bodies.
 - 4. A method according to claim 3 wherein the ligand molecule is covalently attached to an oil body protein in the oil bodies.
- 5. A method according to claim 2 wherein the ligand is comprised of two molecules, a first molecule that associates with the oil bodies and a second molecule that associates with the target, wherein the first molecule and the second molecule associate with each other.
- 6. A method according to claim 5 wherein the first and second ligand molecules are conjugated to each other.
 - 7. A method according to claim 4 wherein the ligand molecule is a protein.
 - 8. A method according to claim 7 wherein the protein ligand is a fusion protein with the oil body protein.

- 9. A method according to any one of claims 1 to 8 wherein the target molecule is selected from the group consisting of proteins, peptides, organic molecules, lipids, carbohydrates, nucleic acids, cells, cell organelles, cell components, viruses, metals, metal ions and ions.
- 5 10. A method according to claim 8 wherein the ligand molecule is hirudin and the target molecule is thrombin.
 - 11. A method according to claim 8 wherein the ligand molecule is protein A and the target molecule an immunoglobulin.
- 12. A method according to claim 8 wherein the ligand molecule is metallothionein and the target molecule is cadmium.
 - 13. A method according to claim 8 wherein the ligand molecule is a cellulose binding protein and the target molecule is cellulose.
 - 14. A method according to claim 8 wherein the ligand molecule is a nucleic acid binding protein and the target molecule is a nucleic acid.
- 15 15. A method according to claim 14 wherein the ligand is a single stranded DNA binding protein or an RNA binding protein and the target is a single stranded nucleic acid molecule.
 - 16. A method according to claim 2 wherein the ligand is an antibody that binds to the oil body or an oil body protein.
- 20 17. A method according to claim 16 wherein the target is a cell, cell organelle or cell component capable of binding the ligand antibody.

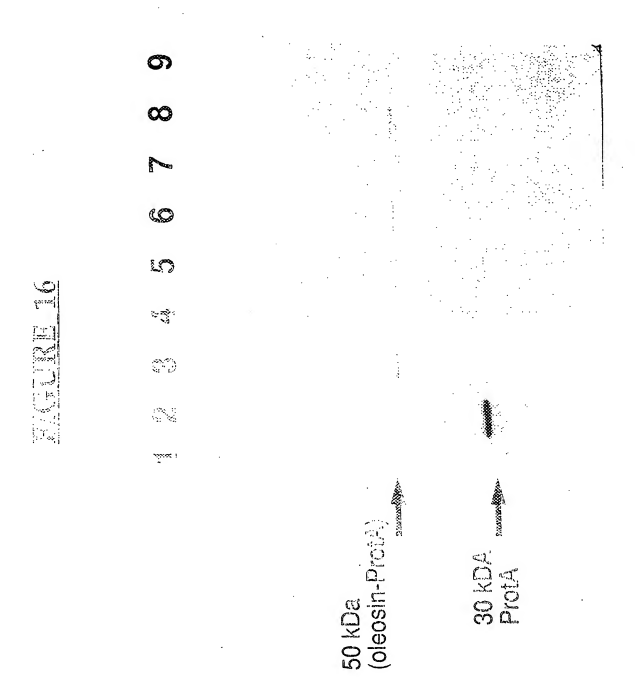
- 18. A method according to claim 16 wherein the cell is Staphylococcus aureus.
- 19. A method according to claim 16 wherein the ligand is a bivalent antibody that binds to both the oil body and the target.
- 5 20. A method according to claim 6 wherein the ligand is an antibody conjugated to avidin and the target molecule is biotin.
 - 21. A method according to any one of claims 4, 7, 8, 9, 10, 11, 12, 13, 14, 15 wherein the oil body protein is an oleosin.
- 22. A method according to claim 21 wherein the oleosin is derived from a plant selected from the group consisting of rapeseed (Brassica spp.), soybean (Glycine max), sunflower (Helianthus annuus), oil palm (Elaeis guineeis), coconut (Cocus nucifera), castor (Ricinus communis), safflower (Carthamus tinctorius), mustard (Brassica spp. and Sinapis alba), coriander (Coriandrum sativum) linseed/flax (Linum usitatissimum), thale cress (Arabidopsis thaliana) and maize (Zea mays).
 - 23. A method according to any one of claims 1 to 22 wherein in step 1) the oil bodies and the sample are mixed and then incubated for about 1 minute to about 24 hours.
- A method according to claim 23 wherein the mixed oil bodies and sample are incubated at a temperature range from about 4°C to about room temperature.
 - A method according to any one of claims 1 to 24 wherein the oil bodies associated with the target molecule are separated from the sample in step (2) by centrifugation, floatation or size exclusion.

- 26. A method according to any one of claims 1 to 25, further comprising 3) separating the target molecule from the oil bodies.
- 27. A method according to claim 26 wherein the target molecule is separated by elution under appropriate conditions.
- 5 28. A method according to any one of claims 1 to 27 wherein the oil bodies are obtained from the group of plants consisting of rapeseed (Brassica spp.), soybean (Glycine max), sunflower (Helianthus annuus), oil palm (Elaeis guineeis), coconut (Cocus nucifera), castor (Ricinus communis), safflower (Carthamus tinctorius), mustard (Brassica spp. and Sinapis alba), coriander (Coriandrum sativum) linseed/flax (Linum usitatissimum), thale cress (Arabidopsis thaliana) and maize (Zea mays).
 - 29. A composition comprising oil bodies associated with a molecule.
- 30. A composition according to claim 29, wherein the molecule is a target molecule selected from the group consisting of organic molecules, lipids, carbohydrates, nucleic acids, cells, cell organelles, cell components, viruses, metals, metal ions and ions.
- 31. A composition according to claim 30 further comprising a ligand molecule that associates with the oil bodies and the target 20 molecule.
 - 32. A composition according to claim 29, wherein the molecule is a ligand molecule.
 - 33. A composition according to claim 31 or 32 wherein the ligand molecule is covalently attached to the oil bodies.

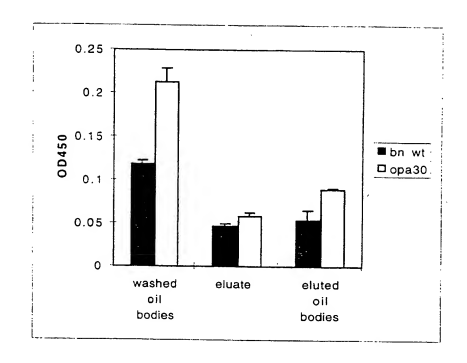
WO 98/27115 PCT/CA97/00951

- 34. A composition according to claim 33 wherein the ligand is biotin.
- 35. An affinity matrix for use in separating a target molecule from a sample comprising an oil body that can associate with the target molecule.
 - An affinity matrix for use in separating a target molecule from a sample comprising (a) an oil body and (b) a ligand molecule associated with the oil body, wherein the ligand molecule is capable of associating with the target molecule.

19/20



26/20 FIGURE 17



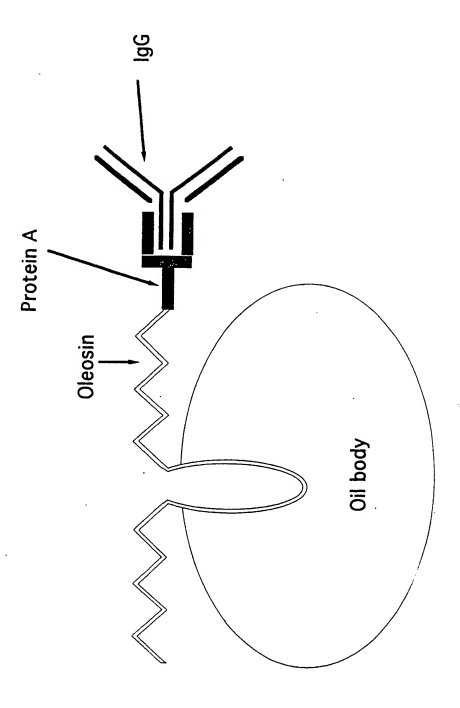


FIGURE 14 CONT'D

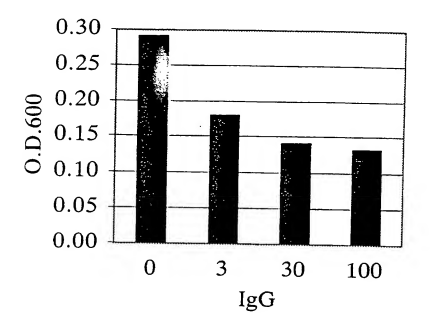
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1504																		-		_	102
163	CGT	GAC	CGT	ACT	CGT	GGT	GGC	CAG									TCC			CAA	165
103	R	D	R	T	R	G	G	Q	н	T	T	<u>L</u>	V	P	R	<u> </u>	s	М_	D	Q	182
1654	CGC	AAT	GGT	ттт	ATC	CAA	AGC	CTT		Cam	Cam		100				AAC				
183	R	N	G	F	I	0	s	L	K	D	D	P	S	O	. AG1	GCT A					171
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1714	GAA	GCT	CAA	AAA	CTT	AAT	GAC	TCT	CAA	GCT	CCA	AAA	GCT	GAT	GCG	CAA	CAA	ААТ	AAC	المنت	177
203	E	A	Q	K	L	N	D	s	Q	A	P	K	A	D	A	Q	Q	N	N	F	222
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303	N	G	F	I	Q	s	L	K	D	D	P	s	Q	S	A	N	L	L	S	E	322
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343	Q	N	A	F	Y	E	I	L.	H	L	P	N	L	N	E	E	Q	R	N	G	362
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474	GAG	CAC	CCA	CAG	GGA	TCA	GAC	AAG	TTG	GAC	AGT	GCA	AGG	ATG	AAG	TTG	4DD		_		
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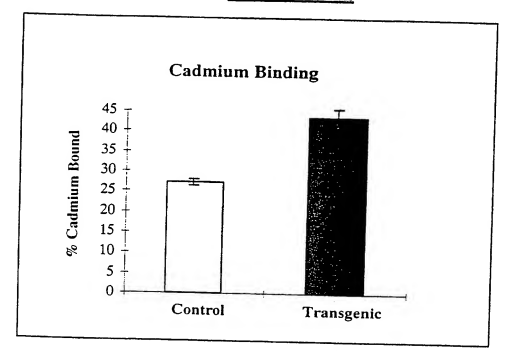
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182	TTA	AAC	GAA	GCG	CAA	CGT	AAC	GGC	TTC	: ATI	CAA	AG:	CTT	KAA 1	A GAC	GAC		A AG	- CA	A AGC	2.
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302	AAT	י איייר	220	222	CAA	CAA	C														
100	N	F	N	K	E	Q	Q	AAT N	GCT	TTC	TAT	GAA	ATC	TTG		ATG	CCI	' AAC	TTA	AAC	36
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362	GAA	GAA	CAA	CGC	AAT	GGT	TTC	ATC	CAA	AGC	тта	222	C N M	C10	CCA						
120	E	E	Q	R	N	G	F	I	0	s	L	K	D	D	P	AGC S					421
									_		_						Q	s	A	N	139
422	CTA	TTG	TCA	GAA	GCT	AAA	AAG	TTA	AAT	GAA	TCT	CAA	GCA	CCG	AAA	GCG	CAT	220	222	mmc.	400
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482	220																				155
160	N	R	E	CAA	CAA	AAT	GCT	TTC	TAT	GAA	ATC	TTA	CAT	TTA	CCT	AAC	TTA	AAC	GAA	GAA	541
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542	CAA	CGC	AAT	GGT	ጥጥር	ATC	CAA	100	Cm.		~										
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02	GCA	GAA	GCT	AAA	AAG	CTA	AAT	GAT	GCT	CAA	GCA	CCA	222	CCT	GAC						
200	A	Ē	A	ĸ	K	L	N	D	A	Q	A	P	K	A		AAC N	AAA K				661
										-								F	N	K	219
62	GAA	CAA	CAA	AAT	GCT	TTC	TAT	GAA	ATT	TTA	CAT	TTA	CCT	AAC	TTA	ACT	GAA	GAA	CAA	CC#	
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22	220																		-		239
40	N	GGC	TTC	ATC	CAA	AGC	CTT	AAA	GAC	GAT	CCG	GGG	AAT	TCC	CGG	GGA	TCC	GTC	GAC	CTG	781
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60	Q				*		dIII														804
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14/20 FIGURE 12

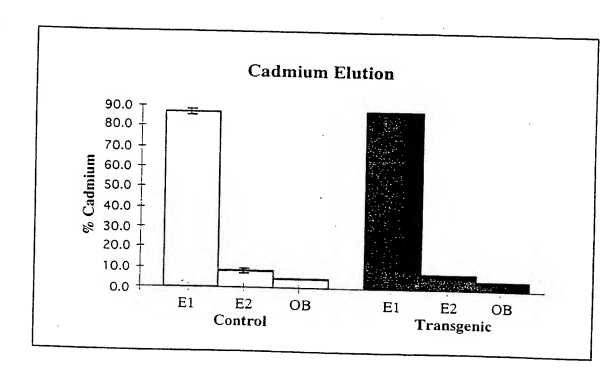


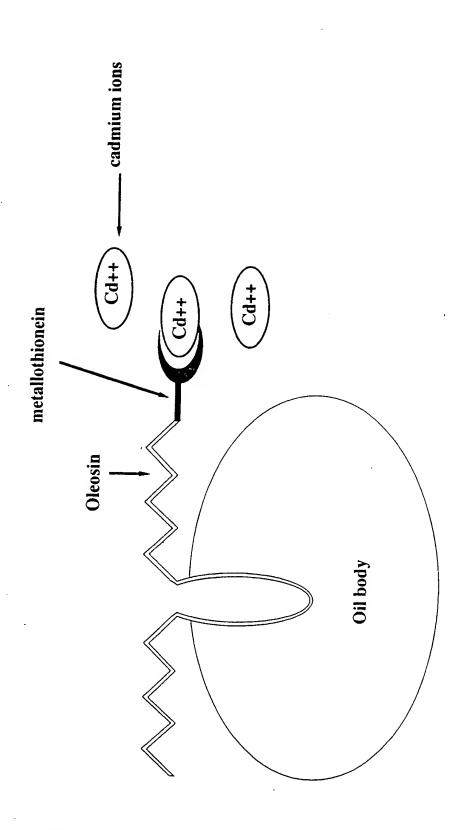
13/20 FIGURE 11

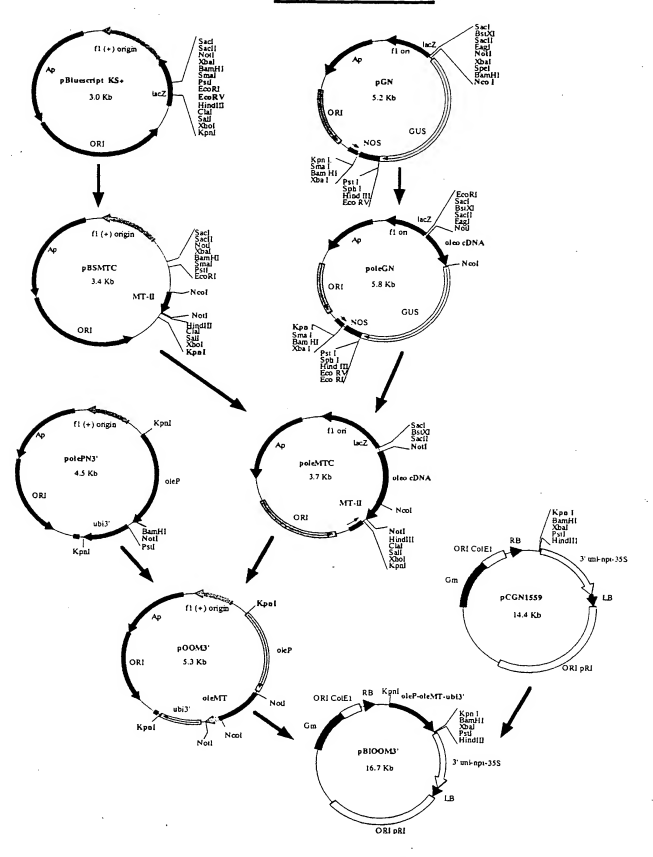
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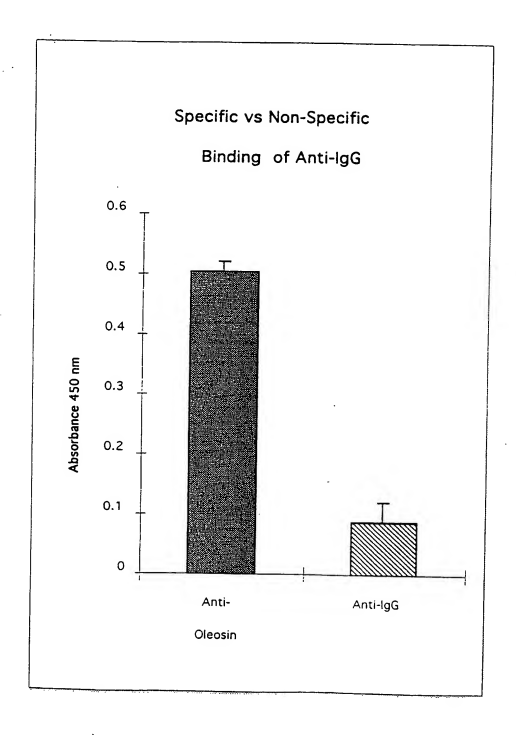
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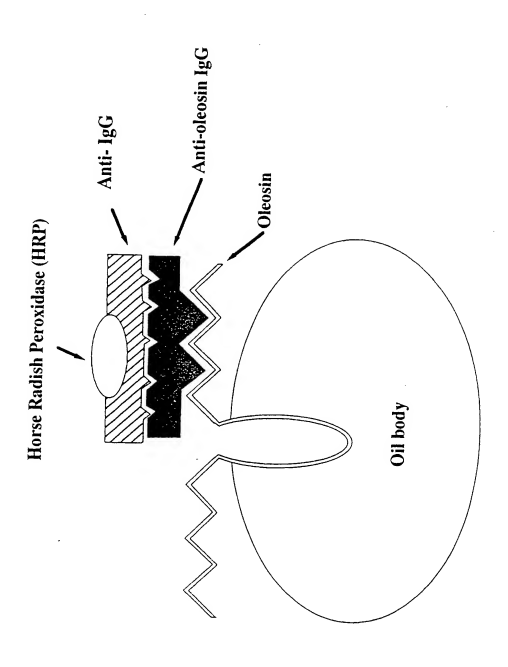
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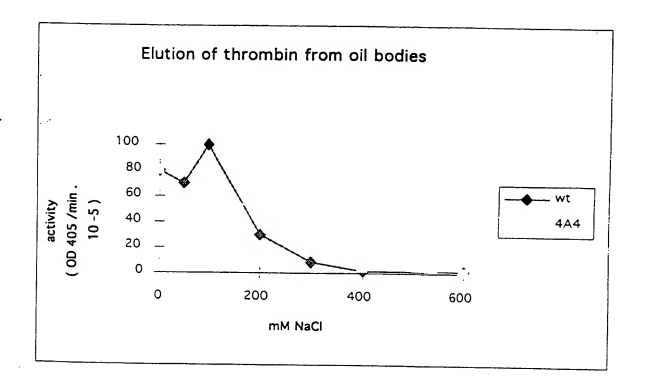
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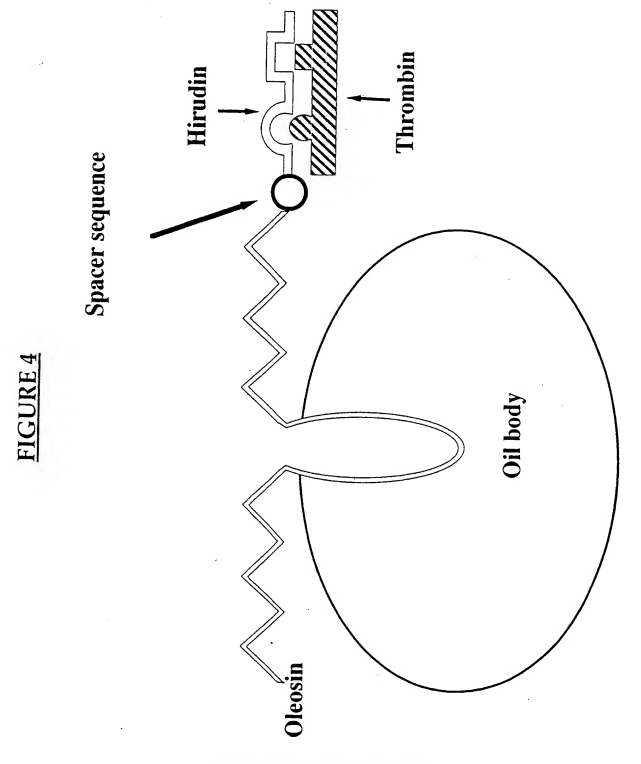
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1833 248	GAC D	AAG K	TGC C	AGC S	TGC C	TGT C	GCC A	TGA •	gcgg	ccgc	gagg	Igctq	caga	atga	gtto	caag	atgg	itttç	gtgac	gaag	1904 255
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1473 128	TCC S	TGG W	ATC I	TAT Y	AAG K	TAC Y	GCA	ACG T	GGA	GAG	CAC	CCA	CAG	GGG	TCA	GAT	AAG	TTG	GAC	AGT	1532
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148	A	R	М	ĸ	L	G	T	AAA K	GCT A	CAG Q	GAT D	ATT I	AAA K	GAC D	AGA R	GCT A	CAA	TAC		GGA	1592
1593 168	CAG	CAA	CAT	ACA	GGT	GGT	GAG	CAT	GAC	CGT	GAC	CGT	ACT	ርርጥ	GCT	ece.	~~~		Y	G	167
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188	L	GTT V	CCA P	CGA R	GGA G	TCC S	ATG M	GAT D	CCC	AAC	TGC	TCC	TGT	GCC	GCC	AGT	GAC	TCC	TGC	ACC	1712
						_	-	_	-	47	С	S	С	A	A	S	D	s	С	T	207



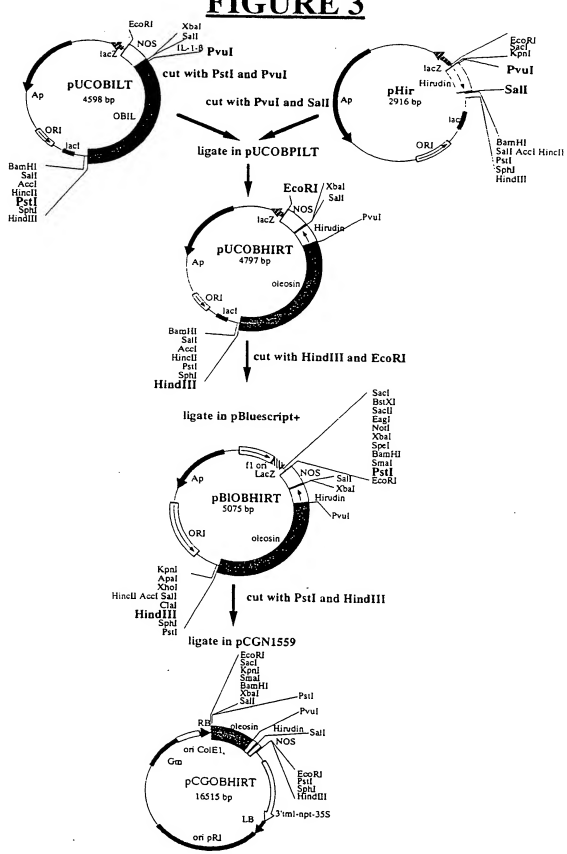






SUBSTITUTE SHEET (RULE 26)

FIGURE 3



SUBSTITUTE SHEET (RULE 26)

3/20 FIGURE 2 CONT'D

1651 184	TGT C	ACT T	GAA E	TCT S	GGA G	CAG Q	AAC N	CTC L	TGT C	CTC L	TGT C	GAA E	GGA G	TCT S	AAC N	GTT V	TGT C		aag K	GGA G	171 203
1711 204	AAC N	AAG K	TGT C	ATC I	CTC L	GGA G	TCT S	AAC N	GGA G	aag K	GGA G	AAC N	CAG Q	TGT C	GTT V	ACT T		GAA E	GGA G	ACT T	1770 223
1771 224	CCA P	AAC N	CCA P	GAA E	TCT S	CAC H	AAC N	AAC N	GGA G	GAC D	TTC F	GAA E	GAA E	ATC I	CCT P				_	CAG O	1830 243
1831 -244	TAA •	gtcg	acto	taga	acgga	tctc	ccga	tegt	tcaa	acat	ttgg	caat	aaag	ittc	ttaa	ıgatt	gaat	.cctg	ittgc	cggt	1909
1910	cttg	cgat	gatt	atca	tata	attt	ctgt	tgaa	ttac	gtta	agca	tgta	ataa	ttaa	cato	taat	acat			ttat	
1990	gaga	tggg	ttt	tatg	atta	gagt	cccg	caat	tata	catt	taat	acac	cata	caaa	2022			gacy	ccac	tagg	1989
2070	ataa	atta	tcac	acac	aata	tcat	ctat	 -					guca	yaaa	acaa	aata	tage	gcgc	aaac	tagg	2069
			, -	3-3-	22,03		cuac	ycta	ctag	accu	GAAT	TC									2115

																				gccaa	
81	ttq	accd	gaatt	gct	gacc	tgaa	gacg	gaac	atca	tcgt	cggg	tcct	tggg	cgat	tgcg	gcgg	aaga	tggg	tcag	cttgg	g 160
161	ctt	gag	gacga	agaco	cga	atcg.	agte	tgtt	gaaa	ggtt	gttc	attg	ggat	ttgt	atac	ggag	attg	gtcg	tcga	gaggt	240
																				gagaga	
321																				tttta	
401																				aatcct	
																				tactga	
																				aaacgt	
																				gcatga	
																				gcatga	
																	TG (
1			•									gcae	acaa	igaac	aaaa	iaa A				ACA T	873 4
874 5	GCT A	AGA R	GGA G	ACC	CAT	CAC	GAT	ATC	ATC	GGC G	AGA	GAC	CAG	TAC					: CG	GAC	933
934	CGA			_			_	_	_	_	R	D	Q	Y	P	М	М	G	R	D	24
25	R	D	Q	Y	Q	M	s	G	R	G	S	D	Y	TCC S	AAG K	TCI S	' AGG	CAC Q	I I	GCT A	993 44
994 45	AAA	GCT A	GCA A	ACT T	GCT	GTC	ACA	GCT	GGT	GGT	TCC	CTC	CTT	GTT	CTC	TCC	AGO	CTT	ACC	CTT	1053
•••	•	^	^	1	A	V	T	A	G	G	s	L	L	V	L	s	S	L	T	L	64
65	V	GGA	ACT T	GTC V	ATA I	GCT A	TTG L	ACT T	GTT V	GCA A	ACA T	CCT P	CTG L	CTC L	GTT V	ATC I	TTC F	AGC S	CCA	ATC	1113 84
1114	CTT	GTC	CCG	GCT	CTC	ATC	ACA	GTT	GCA	CTC	CTC	ATC	ACC	GGT	TTT	СТТ	TCC	TOT	GGA	-	1173
	_	•	F	A	L	1	T	V	A	L	L ·	I	T	G	F	L	s	s	G	G	104
1174 105	TTT F	GGC G	ATT I	GCC A	GCT A	ATA I	ACC T	GTT V	TTC F	TCT S	TGG W	ATT I	TAC Y	AA K	gtaa	gcac	acat	ttat	catc	ttact	
1240	tca	taati	tttgi	tgca	atat	gtge	atgo	atgt	atta	agcc	agta	_	_							atgta	118
																				atgta tctgt	
																					1399
1400 119					-990		cycc	-gg c	icga	cace	catt	gatt	grgaa	atag	G T	AC GO A	CA AC T	CG G G	GAG. E	AG	1470 123
1471 124	CAC	CCA P	CAG Q	GGA G	TCA S	GAC D	AAG	TTG	GAC	AGT	GCA	AGG	ATG	AAG	TTG	GGA	AGC	AAA	GCT	CAG	1530
		-	•	•	•			L	ט	S	А	R	М	K	L	G	S	K	A	Q	143
1531 144	D	L	K K	GAC D	AGA R	GCT A	CAG Q	TAC Y	TAC Y	GGA G	CAG Q	CAA Q	CAT H	ACT T	GGT G	TGG W	GAA E	CAT H	GAC D	CGT R	1590 163
1591 164	GAC	CGT	ACT	CGT	GGT	GGC	CAG	CAC	ACT	ACT	GCG	ATC	GAA	GGG	AGA	ATC			-		
104	ט	R	T	R	G	G	Q	Н	T	T	A	I	E	G	R	I	T	Y	T	D	1650

	ATG M						•	•		п	U	1	1	G	R	D.	Q	Y	P	м	20
	ATG M						-	•	~	1.1	3	G	R	G	5	D	Y	S	ĸ	S	40
121 41	AGG R	CAG Q	ATT I	GCT A	AAA K	GCT A	GCA A	ACT T	GCT A	GTC V	ACA T	GCT A	GGT G	GGT G	TCC S	CTC L	CTT L	GTT V	CTC L	TCC S	
	AGC S						•	•	•	^	L	1	V	A	T	₽	L	L	v	I	80
	TTC F						_		~	-	1	v	A	Ļ	L	I	T	G	F	L	100
	TCC S									•	•	٧	E	5	W	I	Y	K	Y	A	120
361 [°]	ACG T	GGA G	GAG E	CAC H	CCA P	CAG Q	GGA G	TCA S	GAC D	AAG K	TTG L	GAC D	AGT S	GCA .	AGG .	ATG .	AAG K	TTG L	GGA G		420 140
421 141	AAA K	GCT A	CAG Q	GAT D	CTG .	AAA K	GAC D	AGA R	GCT A	CAG Q	TAC Y	TAC Y	GGA (CAG (CAA (480 160
481 161	CAT H	GAC D	CGT R	GAC D	CGT R	ACT T	CGT R	GGT G	GGC G	CAG Q	CAC H	ACT T		TAA						-	

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INTERNATIONAL SEARCH REPORT

Inter anal Application No PCT/CA 97/00951

A. CLASS IPC 6	SIFICATION OF SUBJECT MATTER		C08B1/00									
According	to International Patent Classification (IPC) or to both national classi	fication and IPC										
B. FIELDS	SEARCHED											
IPC 6		10										
	ation searched other than minimum documentation to the extent tha											
. •	data base consulted during the international search (name of data i	basə and, where practical, search t	erms used)									
	ENTS CONSIDERED TO BE RELEVANT		. –									
Category '	Citation of document, with indication, where appropriate, of the re	elevant passages	Relevant to claim No.									
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A	NL, pages 1167-1180, XP002058961 see the whole document WO 93 21320 A (UNIVERSITY TECHNO INTERNATIONAL INC.) 28 October see the whole document	DLOGIES	1-36									
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	an the priority date claimed	"&" document member of the san										
	actual completion of theinternational search March 1998	Date of mailing of the interna	tional search report									
Name and m	13 March 1998 ne and mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo ni, Fax: (+31-70) 340-3016 Masturzo, P											